

(12) United States Patent

Deshmukh et al.

(54) TISSUE PENETRATION DEVICE

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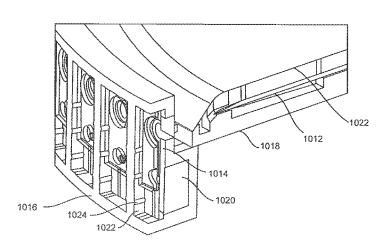
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ABSTRACT

A body fluid sampling system is provided for use on a tissue site. A drive force generator is coupled to penetrating members. The penetrating members are at least partially housed in a penetrating member housing. The penetrating member housing has openings for the penetrating members. The penetrating members are launched along a launch path by the drive force generator through an opening in the penetrating member housing. Analyte sensors are housed in a analyte sensor housing. Each analyte sensor is associated with a penetrating member. The analyte sensor housing and the penetrating member housing are coupled to provide that each analyte sensor is positioned out of a launch path of a penetrating member. The analyte sensor housing includes desiccant pockets for receiving desiccant plugs.

30 Claims, 59 Drawing Sheets



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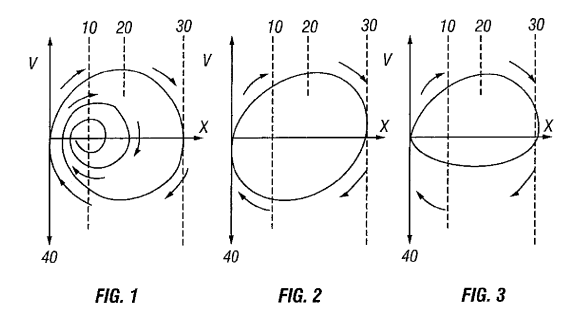
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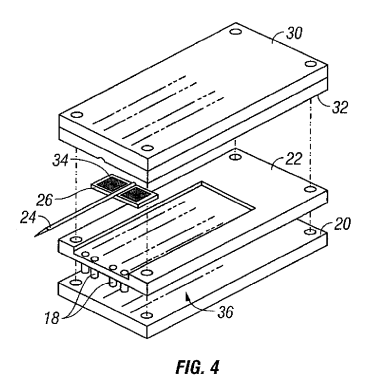
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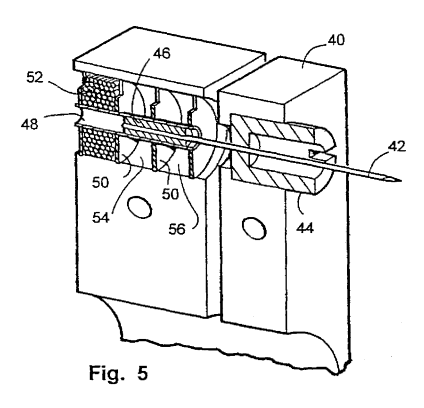
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WO	WO 2006/116441	11/2006	A61B 5/151	69-76)			Cument Comparisons
WO WO	WO 2007/025635 WO 2007/044834	3/2007 4/2007	A61B 5/15 A61B 5/00		tt, W. Heineman, Chrono 20, pp. 121.	coulometry	, Current Separations,
WO WO	WO 2007/054335 WO 2007/070719	5/2007 6/2007	A61B 5/15 A61B 5/00	* cited	1 by examiner		







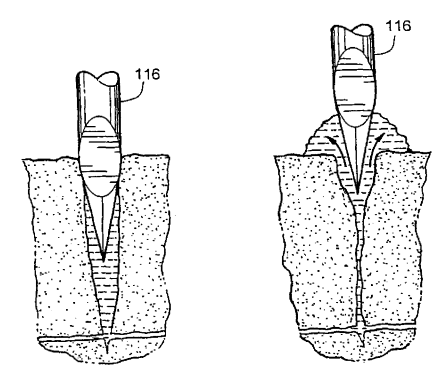
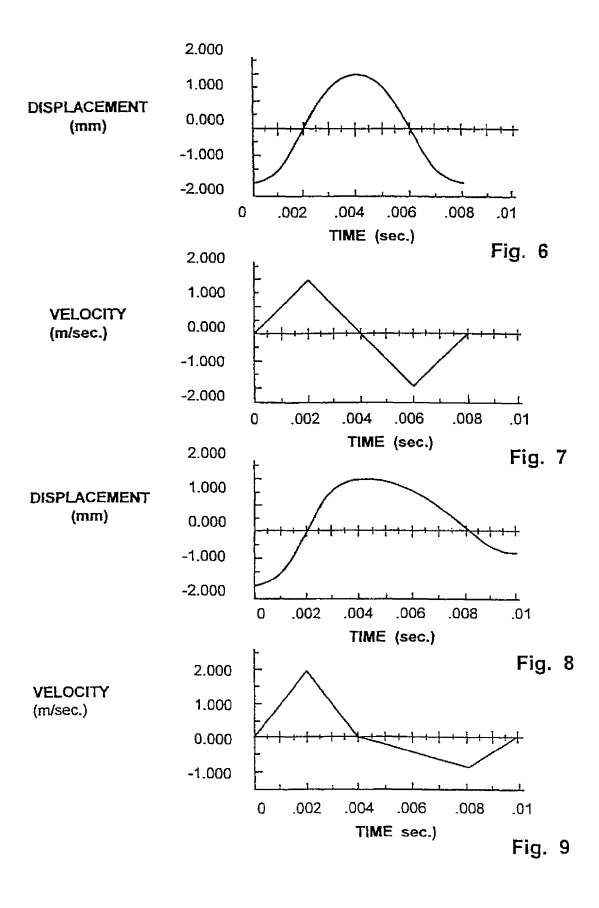


Fig. 10

Fig. 11



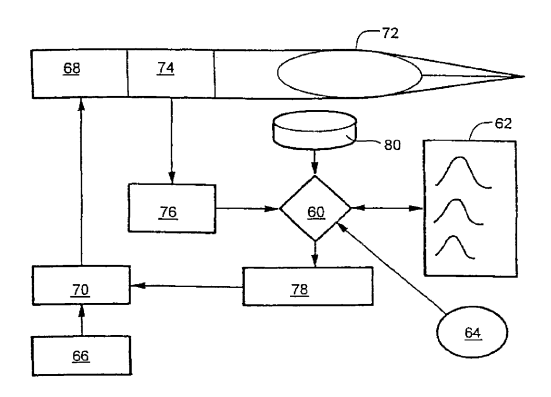


Fig. 12

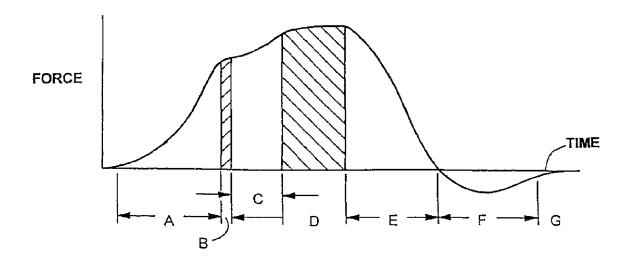
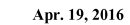


Fig. 13



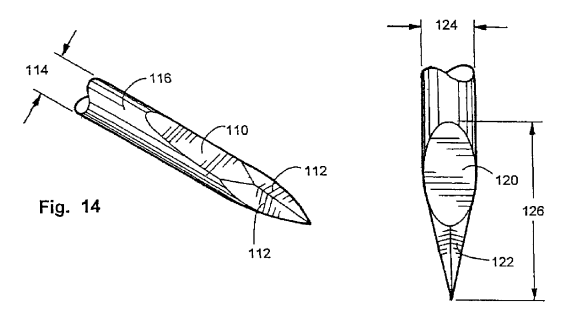
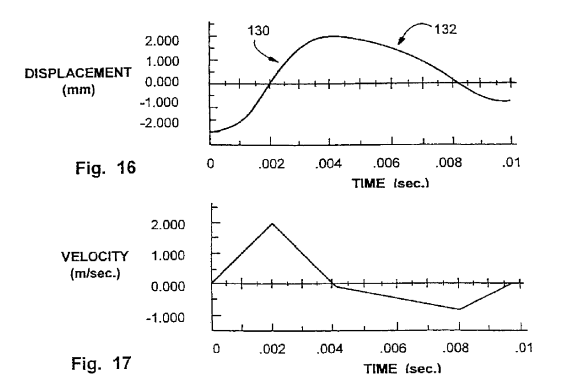


Fig. 15



Apr. 19, 2016

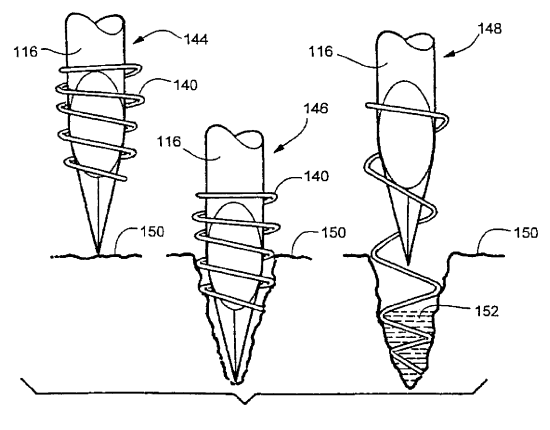


Fig. 18

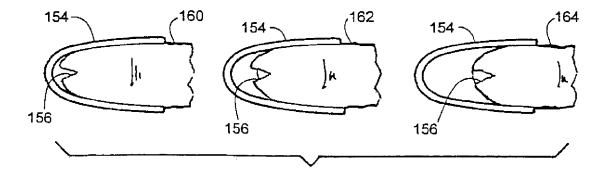
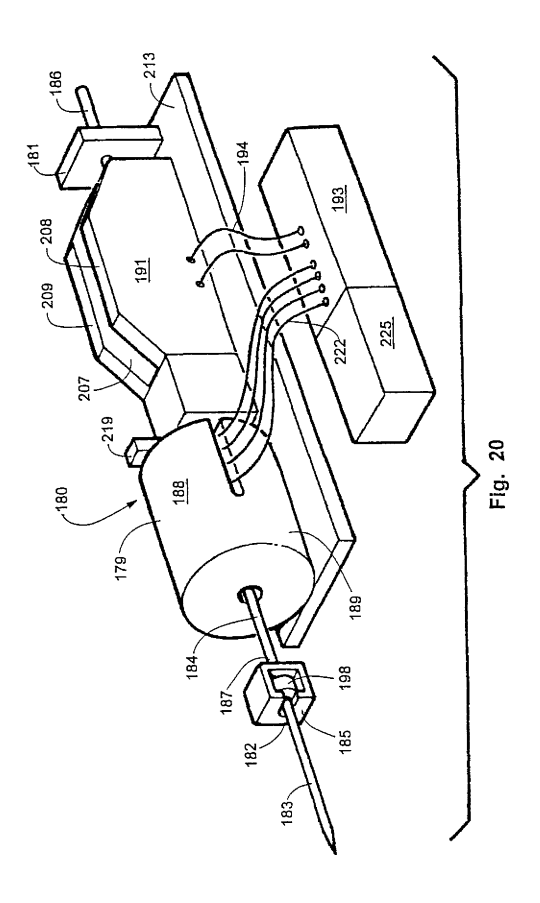
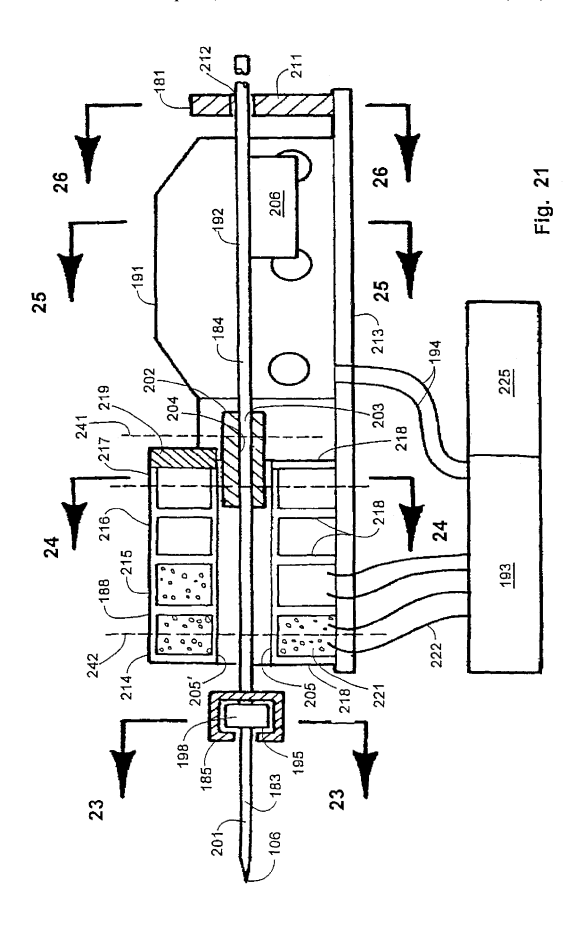


Fig. 19





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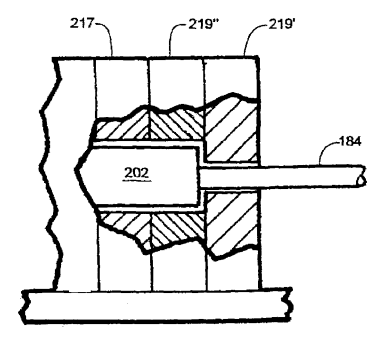


Fig. 22

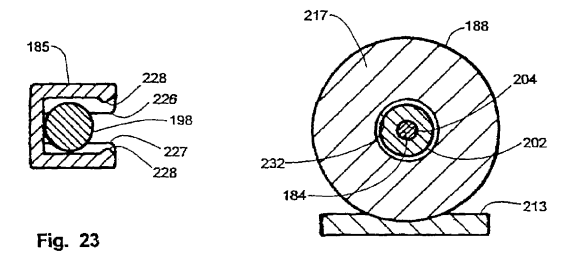
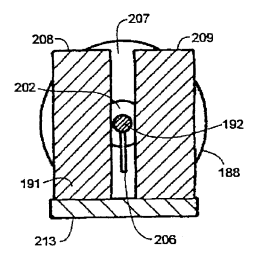


Fig. 24



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Fig. 25

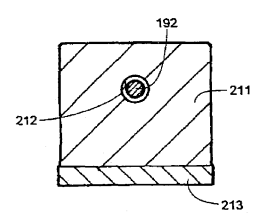


Fig. 26

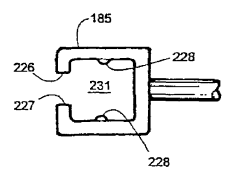


Fig. 27

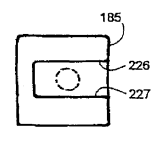
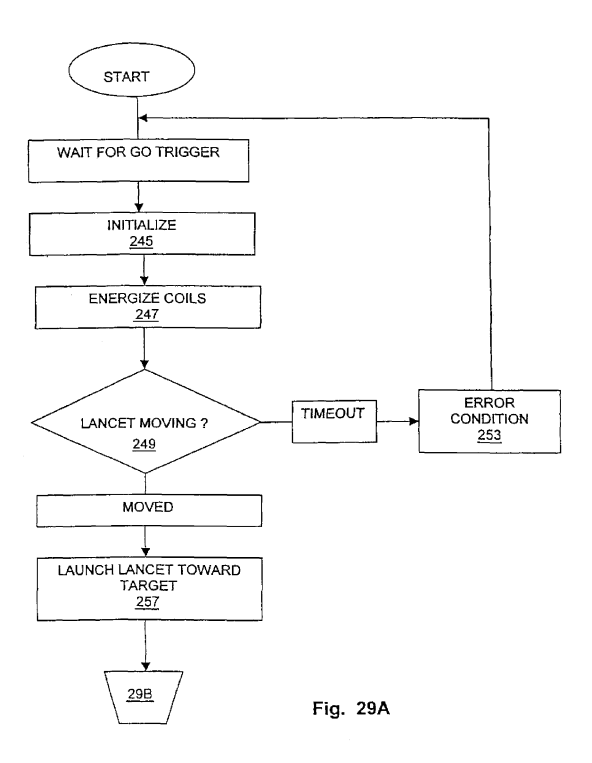


Fig. 28



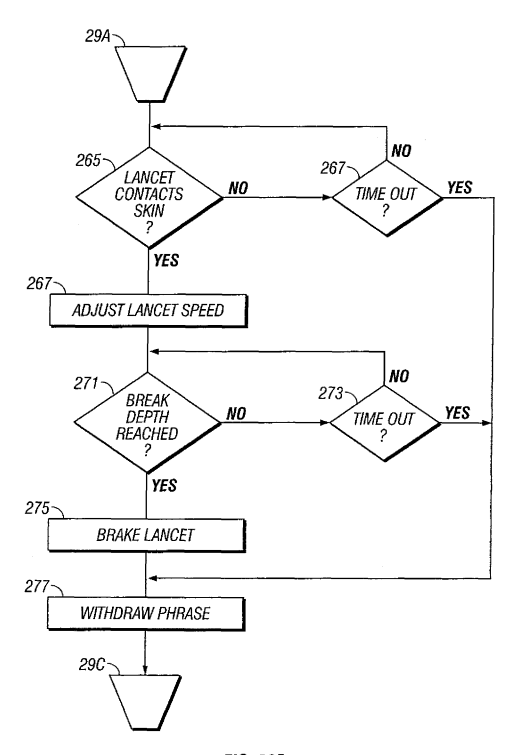
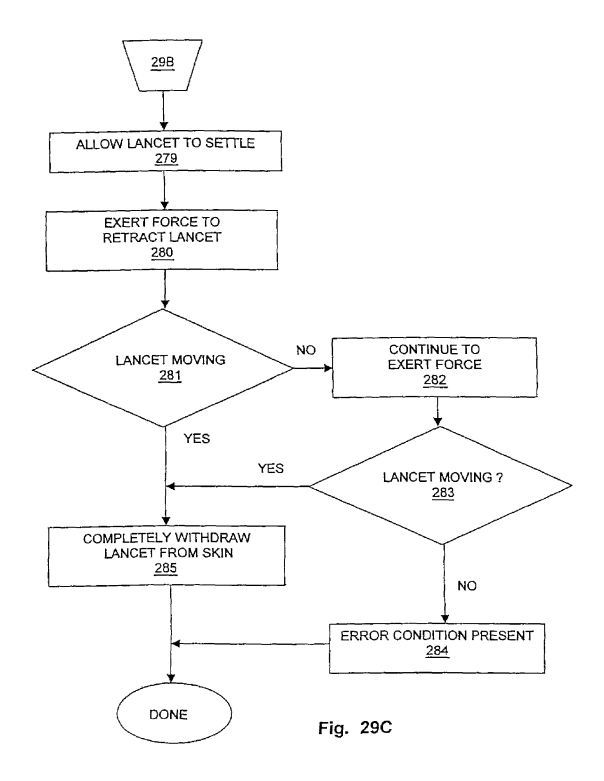
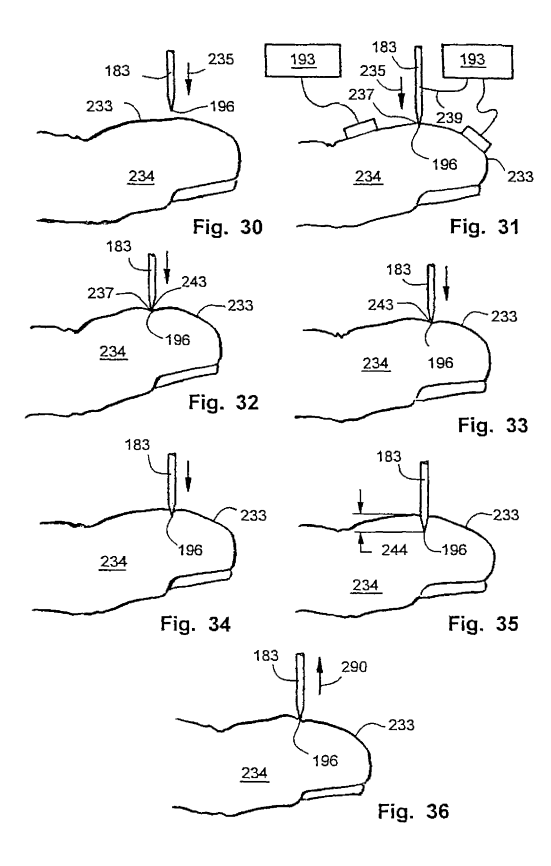
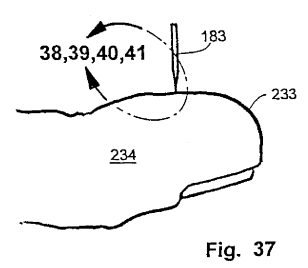


FIG. 29B







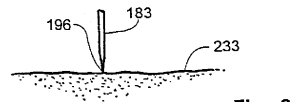
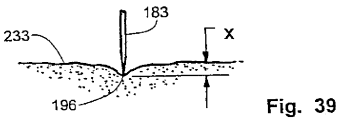


Fig. 38



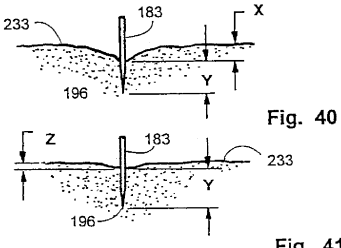
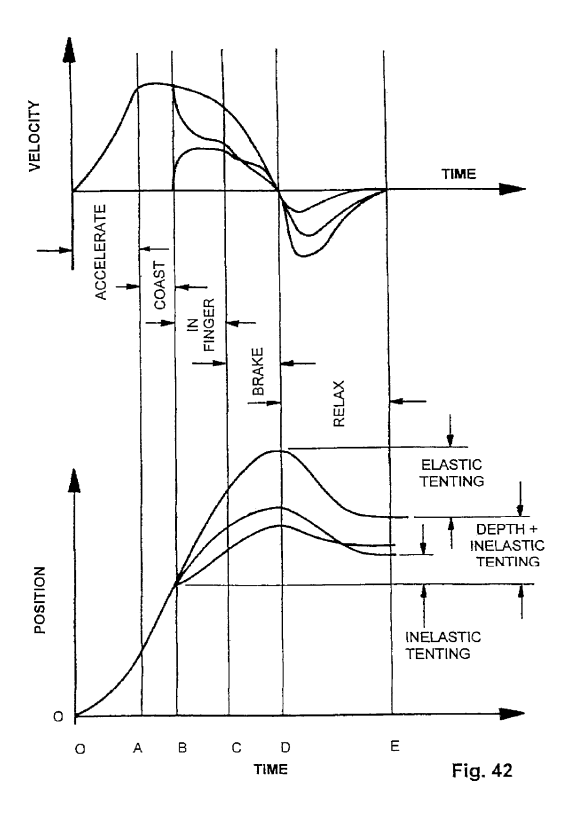
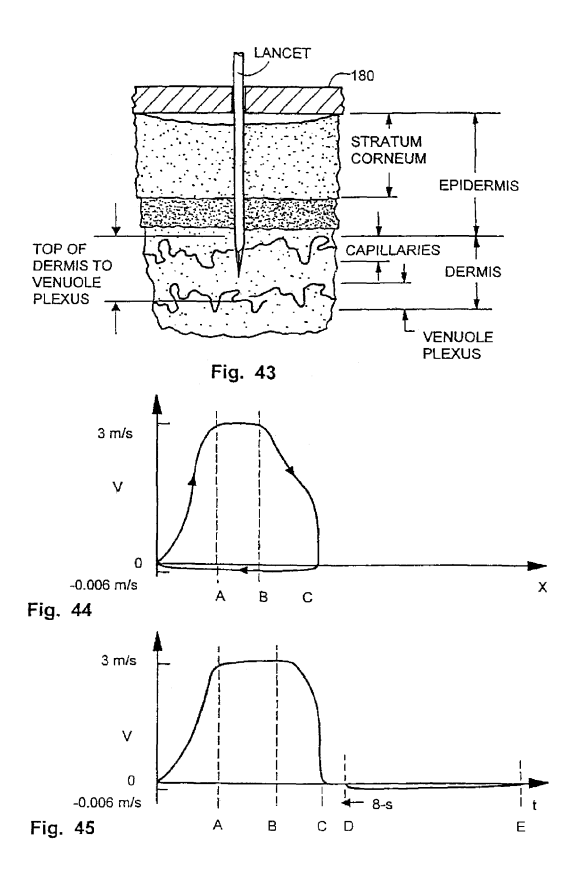
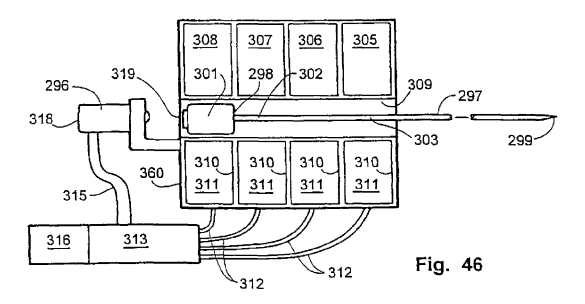


Fig. 41





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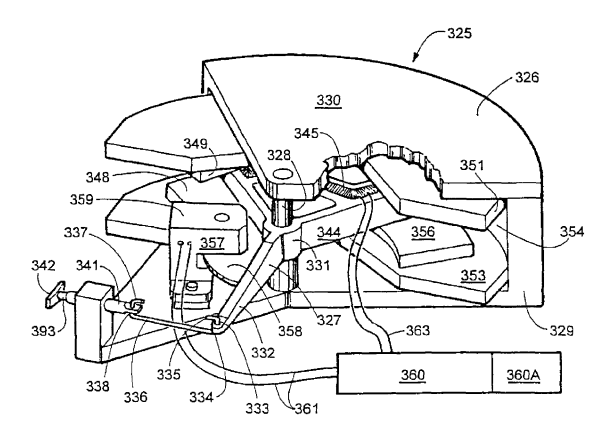
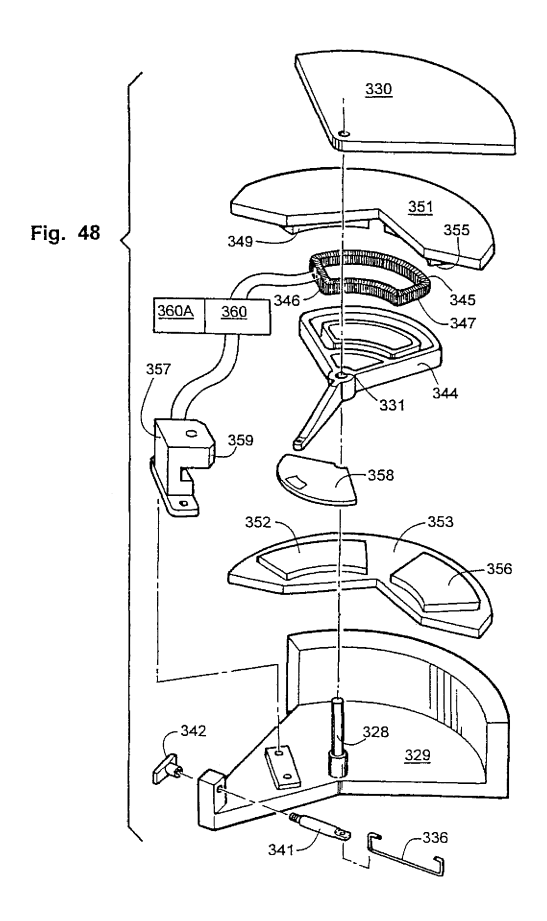
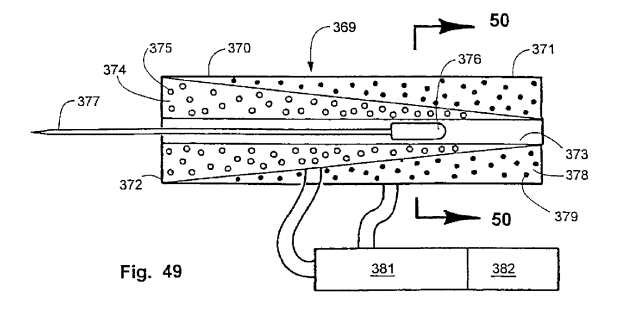


Fig. 47





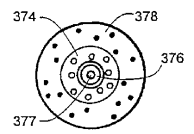
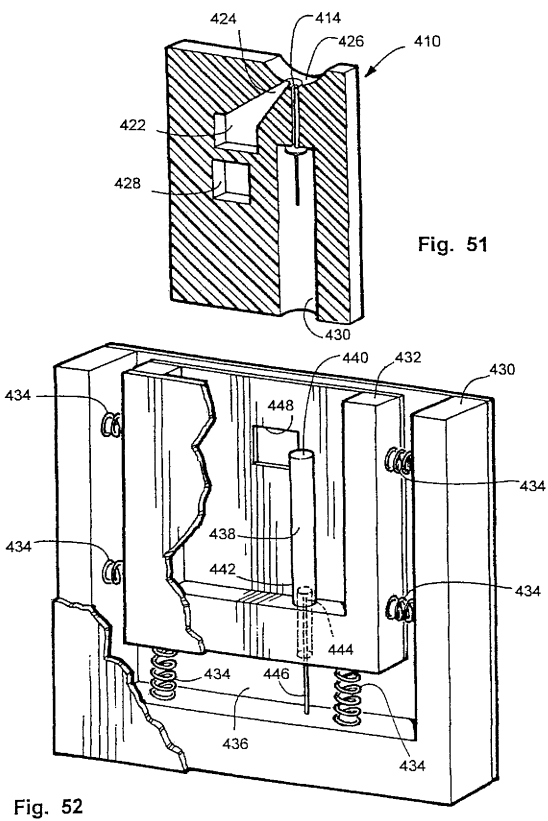


Fig. 50



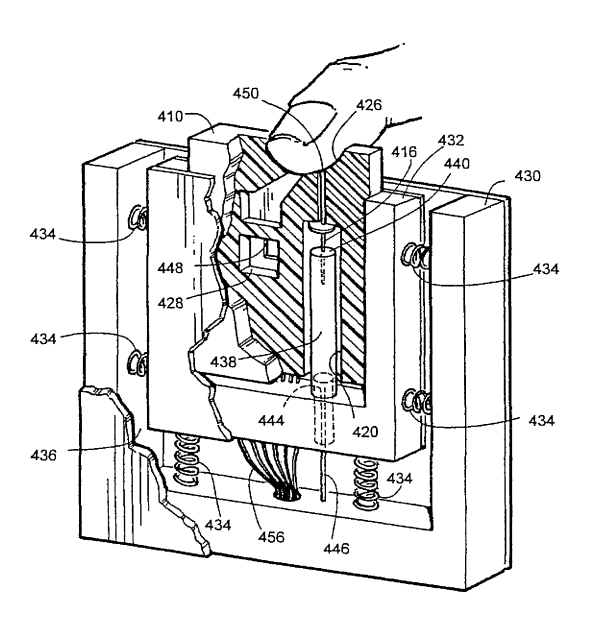


Fig. 53

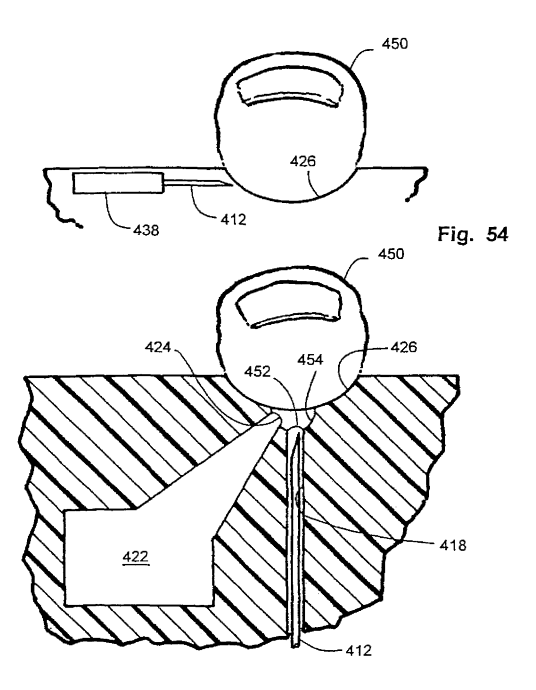


Fig. 55

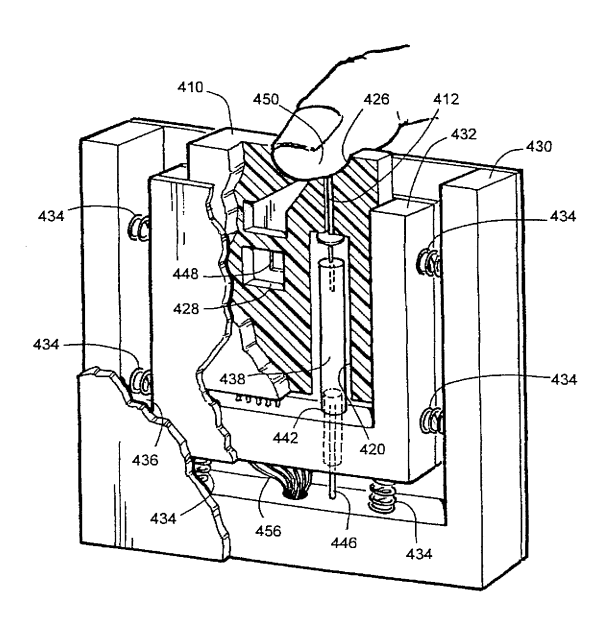


Fig. 56

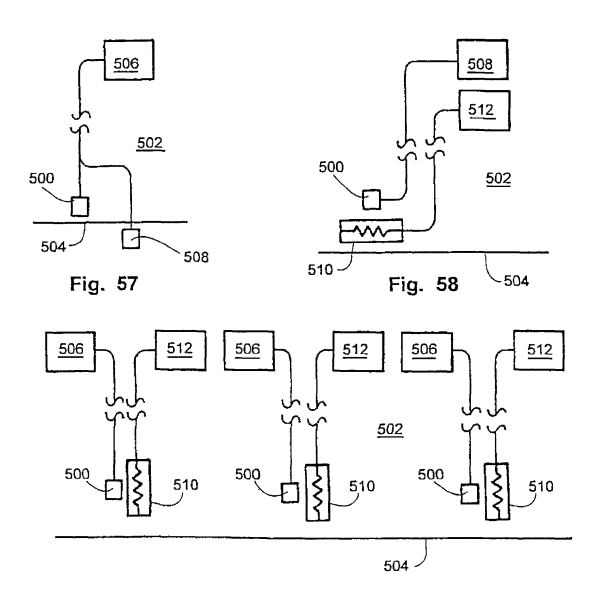
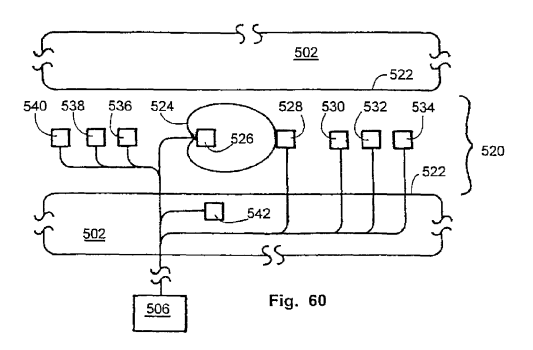


Fig. 59



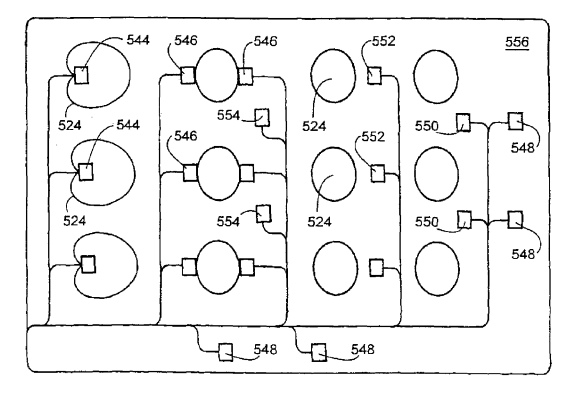
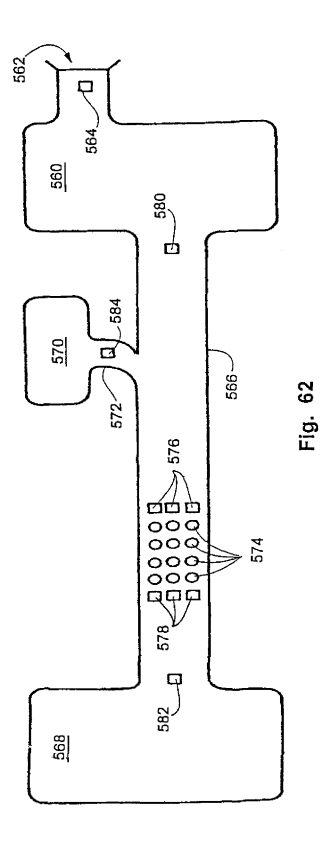
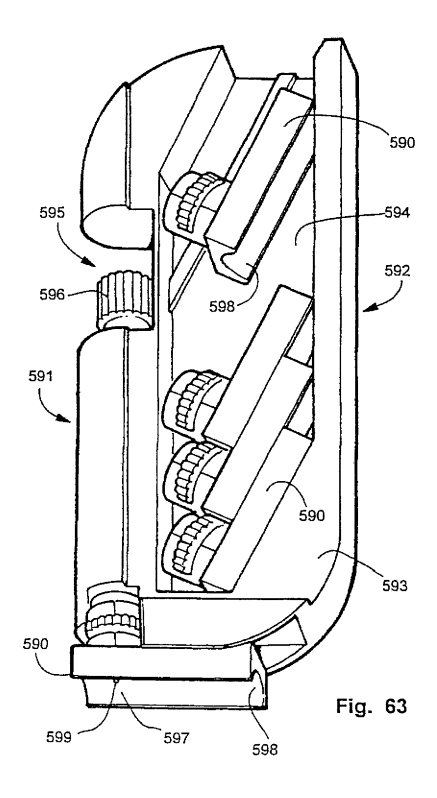
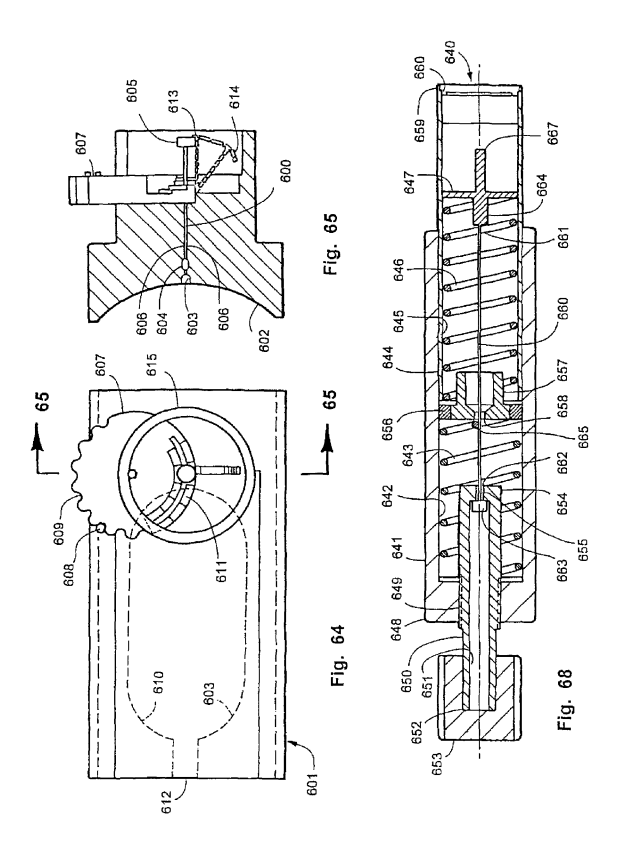


Fig. 61







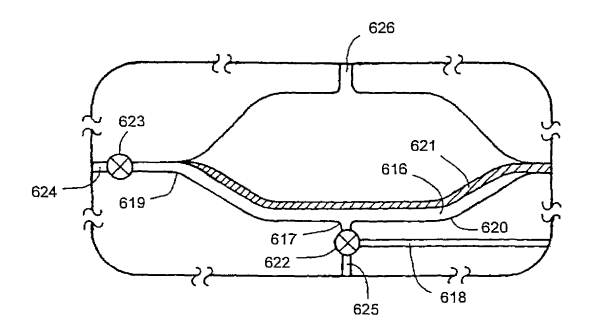
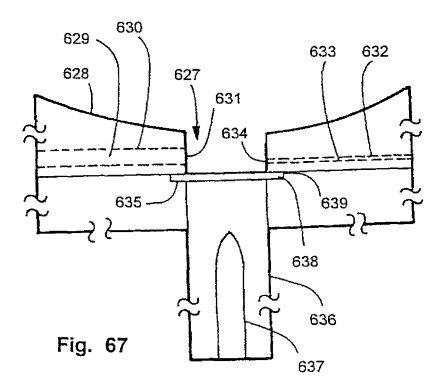
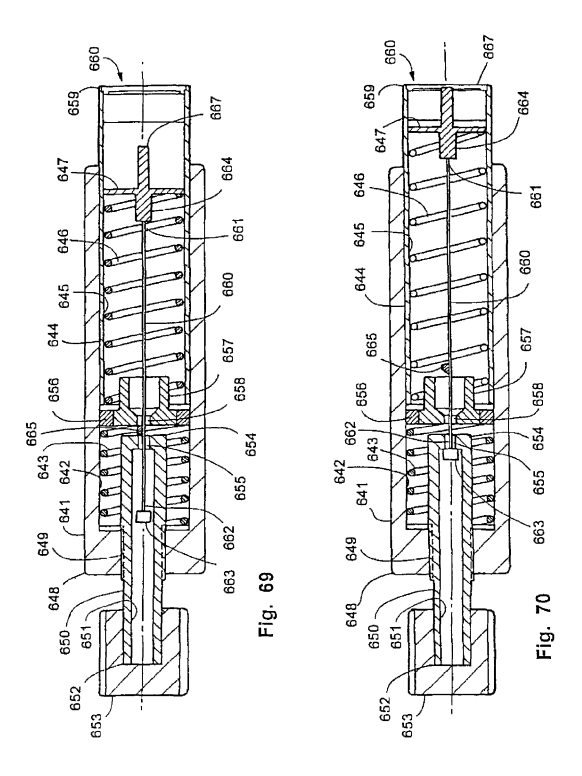
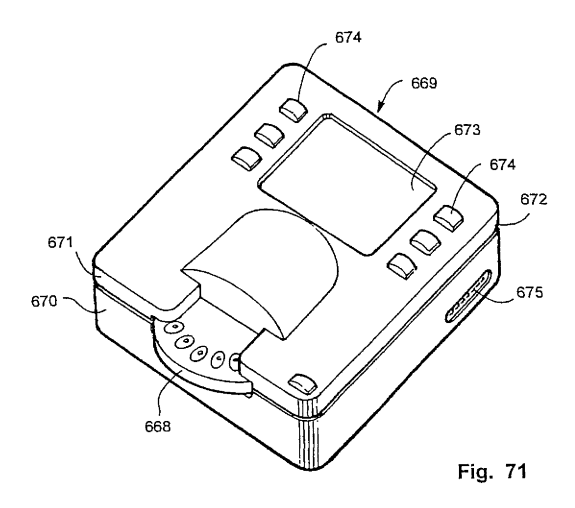
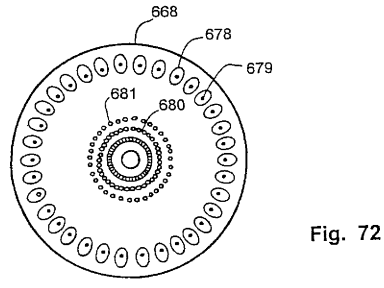


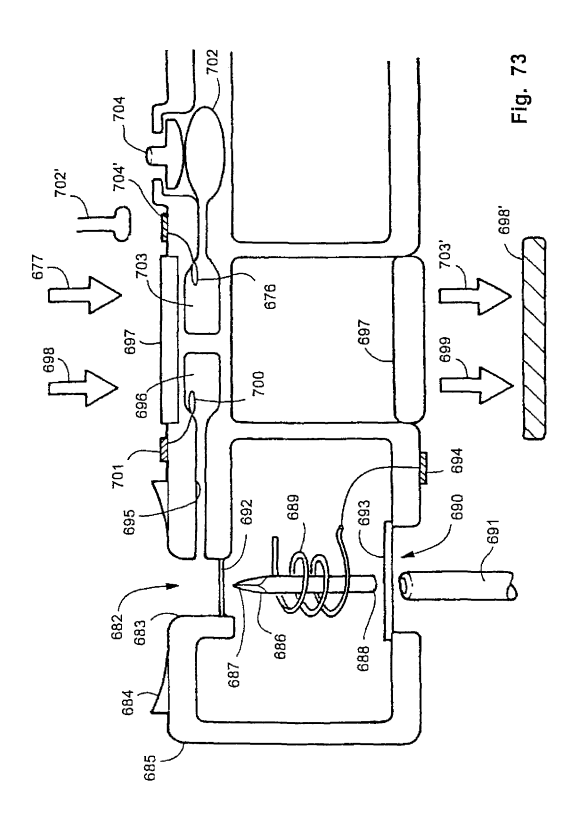
Fig. 66

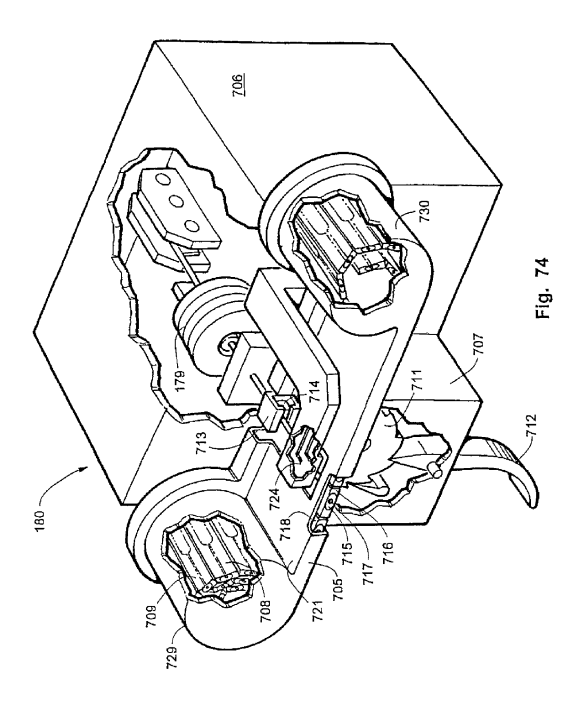


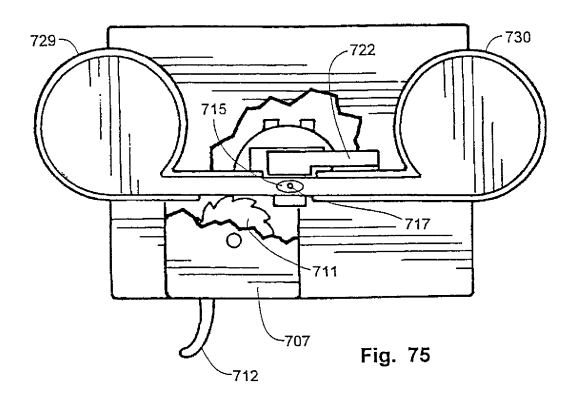












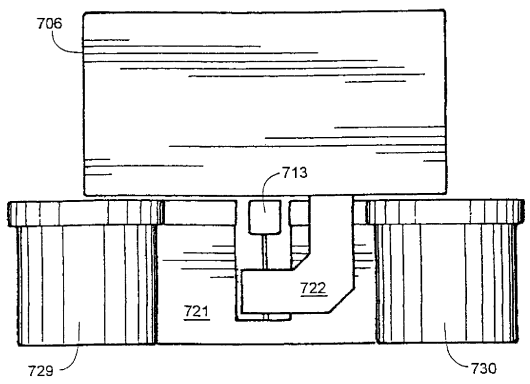
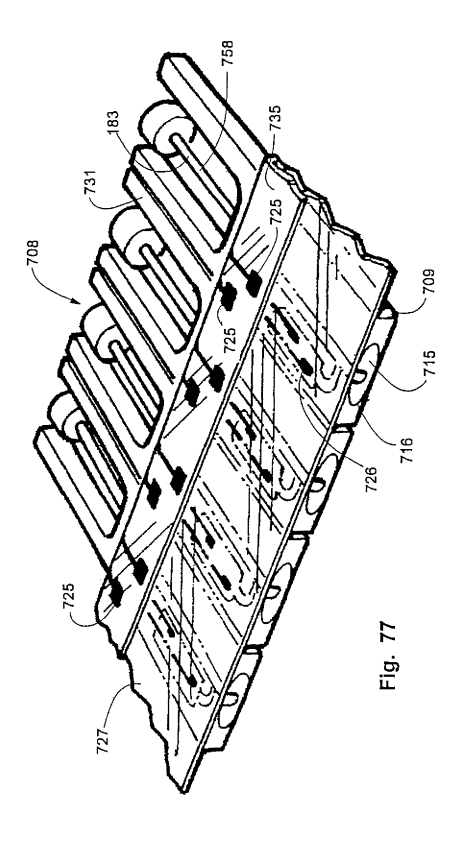
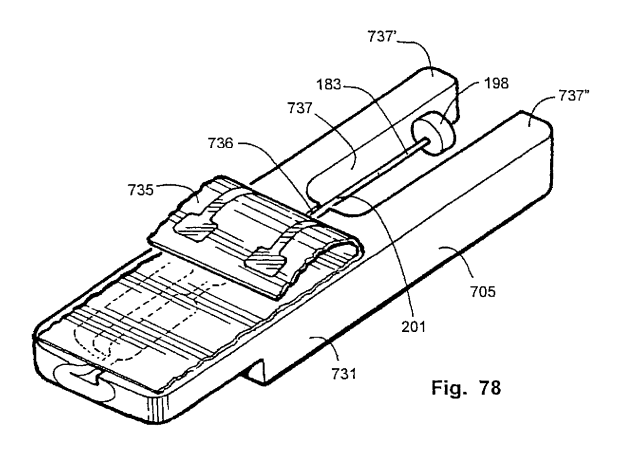
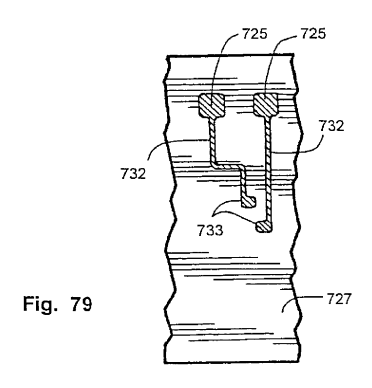
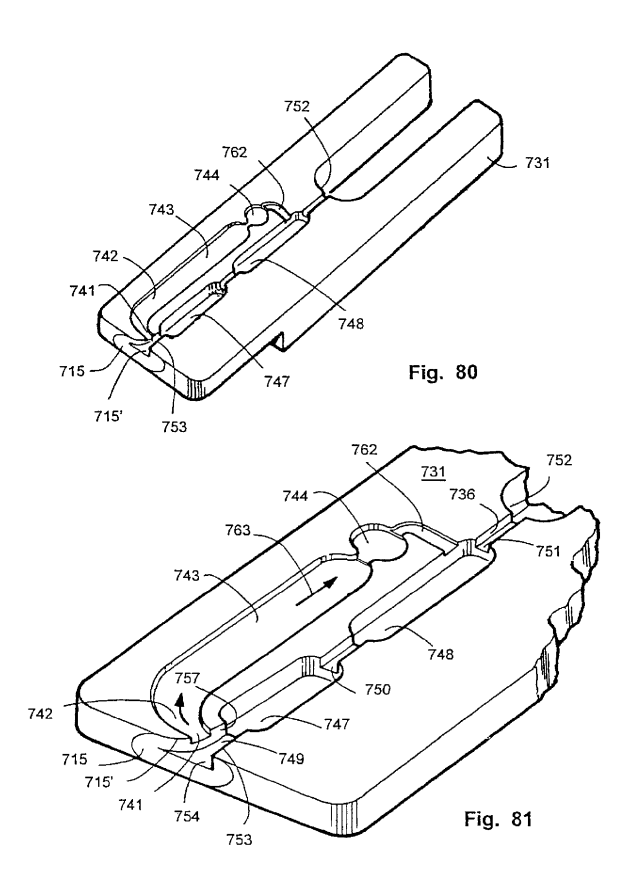


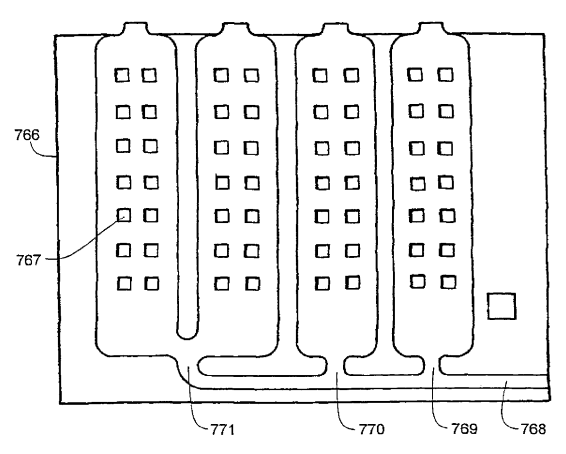
Fig. 76

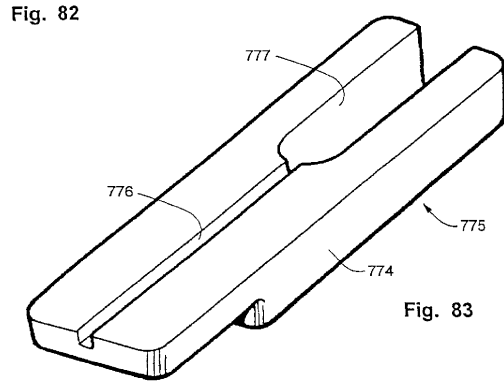


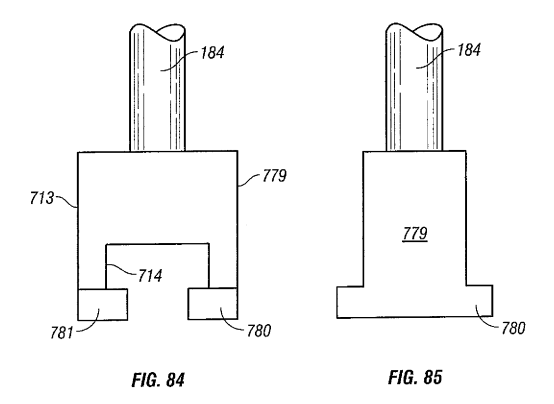


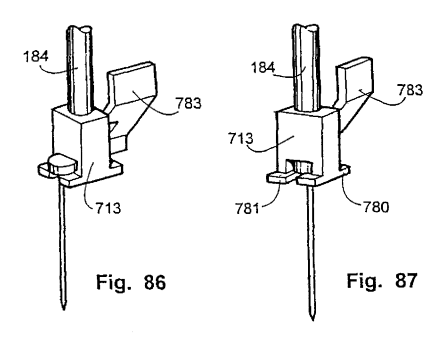


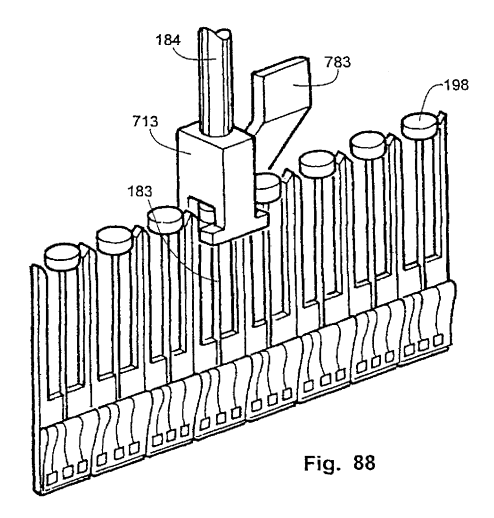


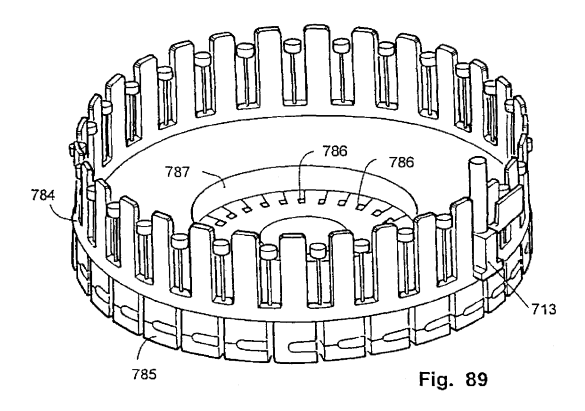


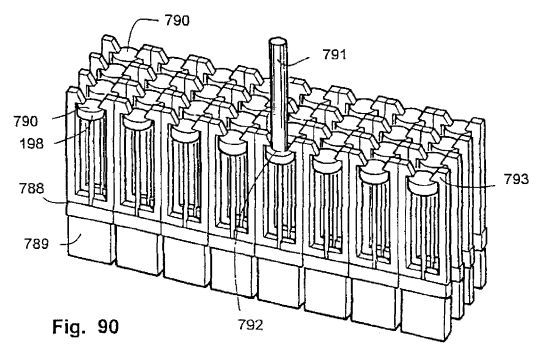


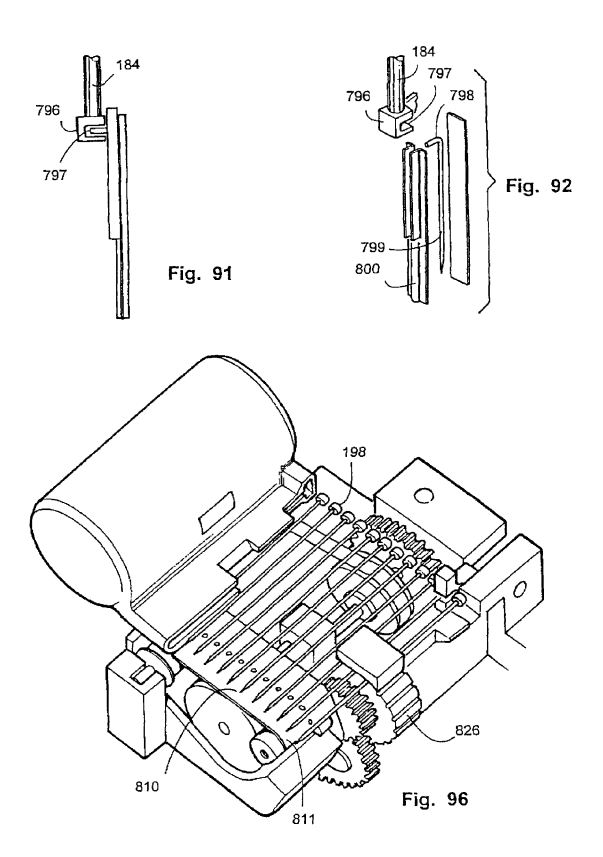


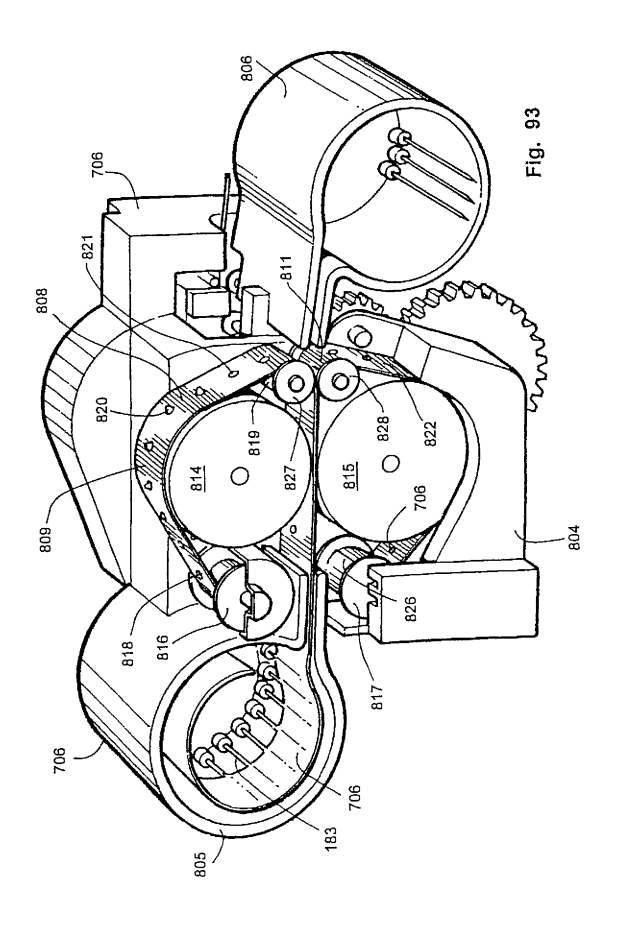


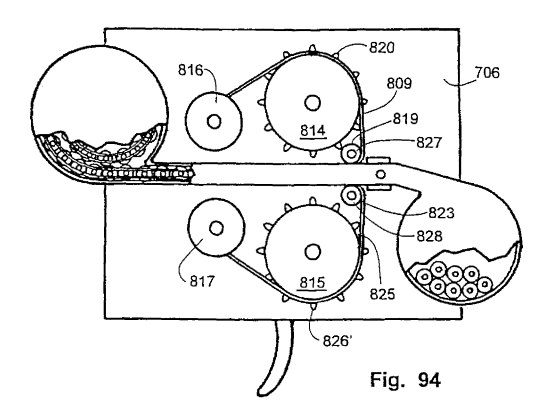


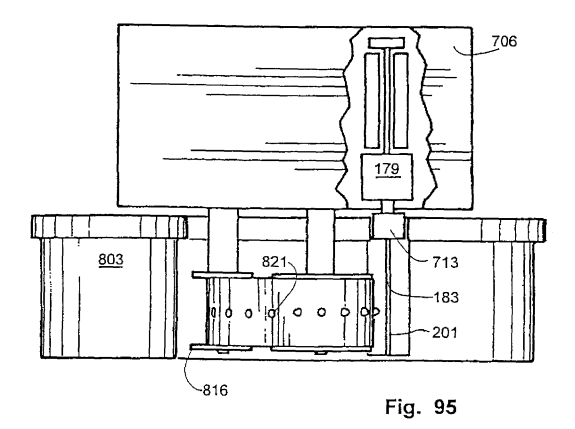


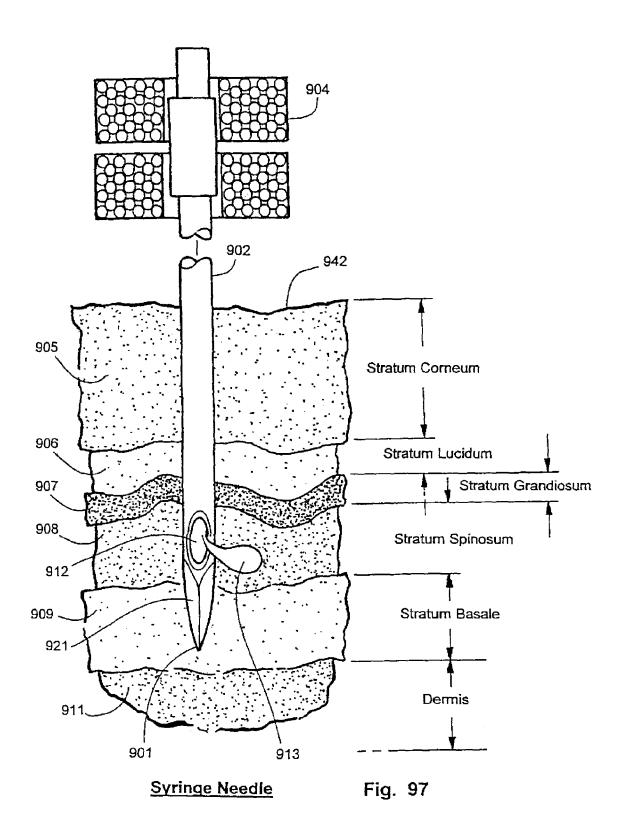


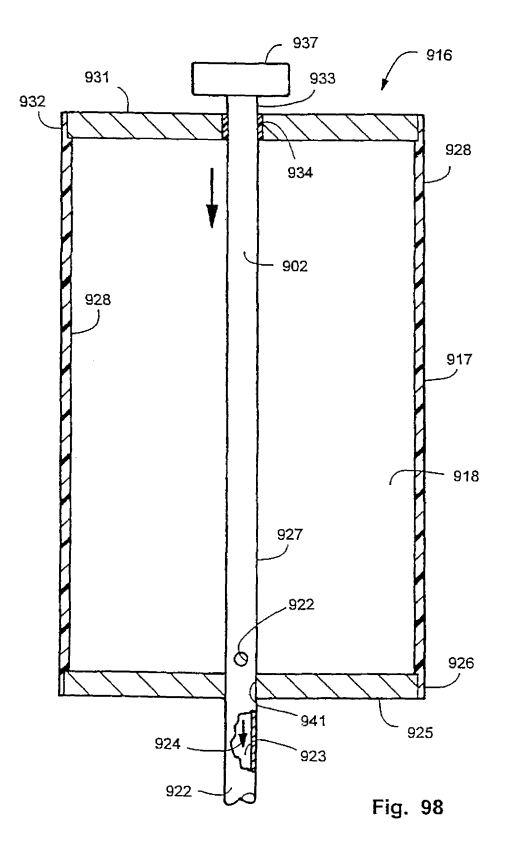


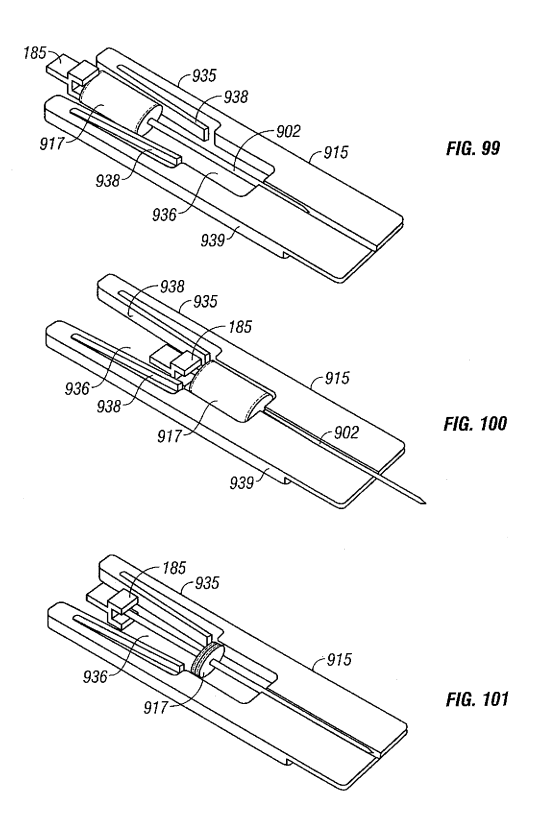


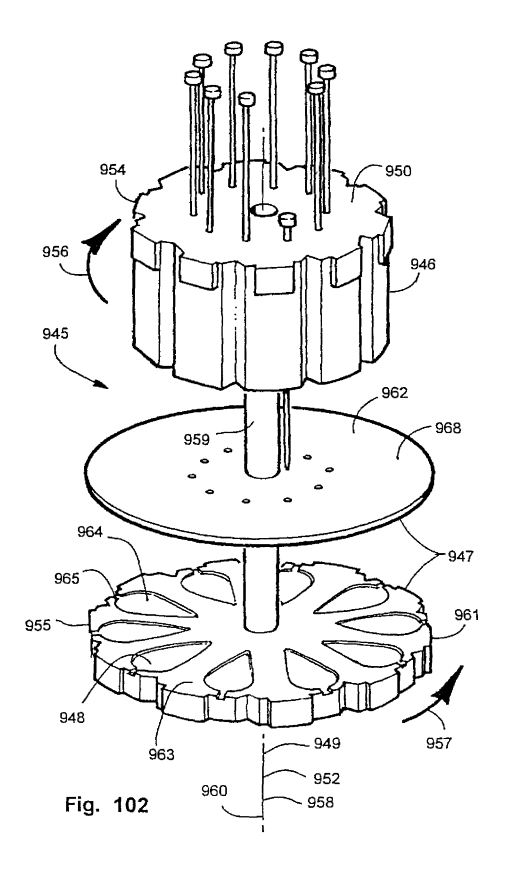


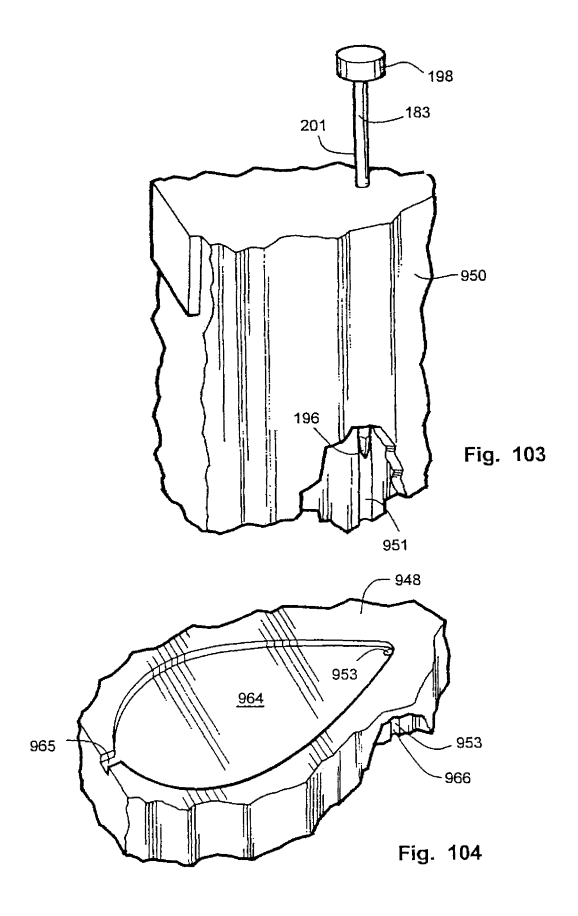


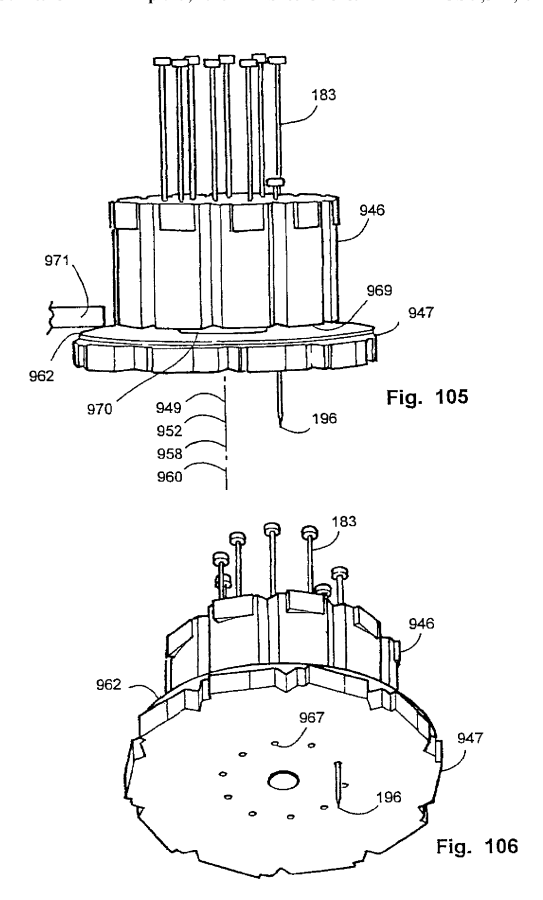


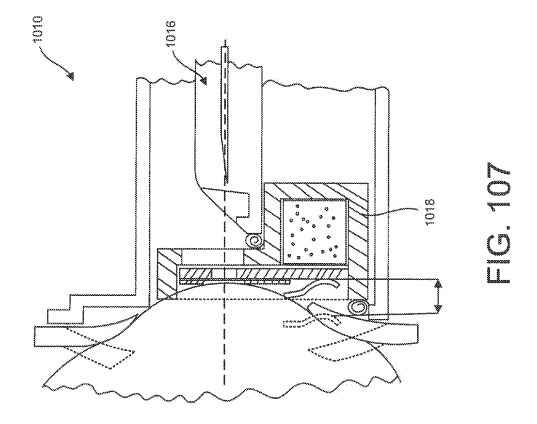


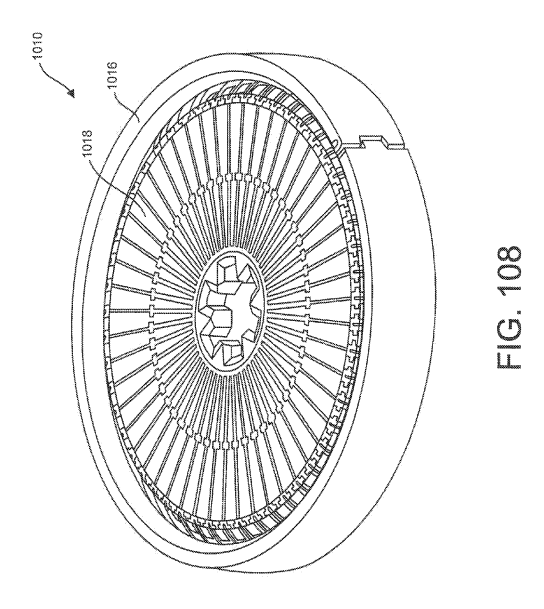


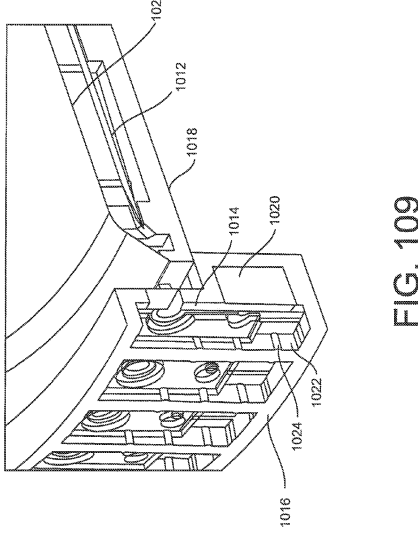












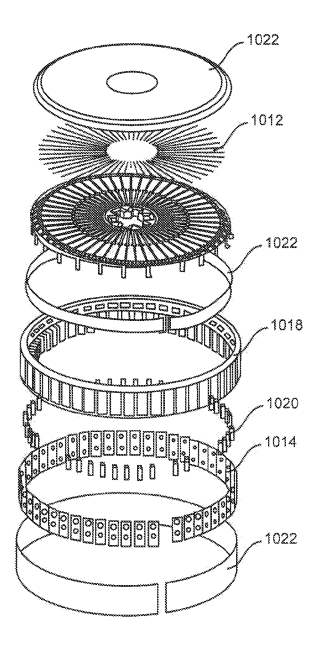
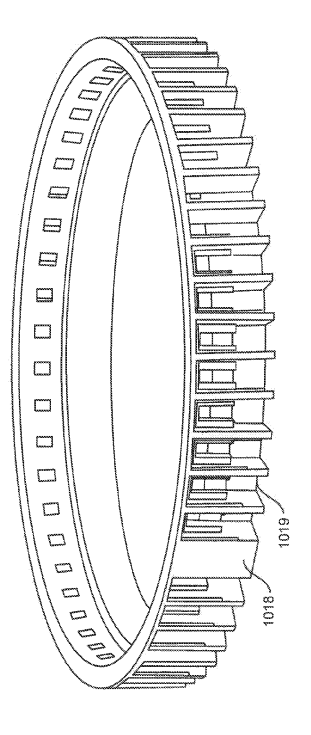
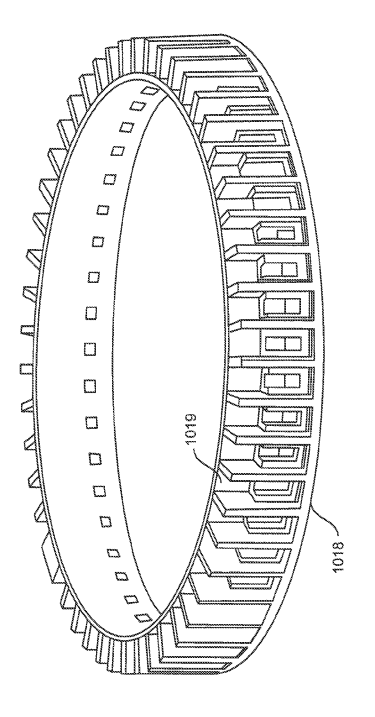
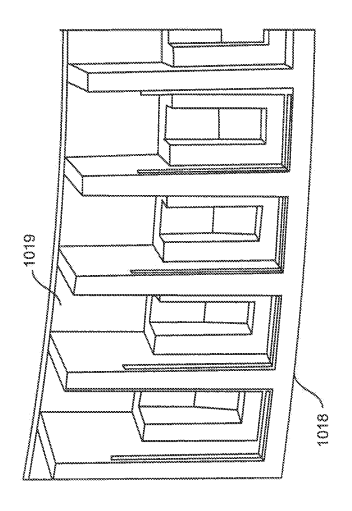


FIG. 110

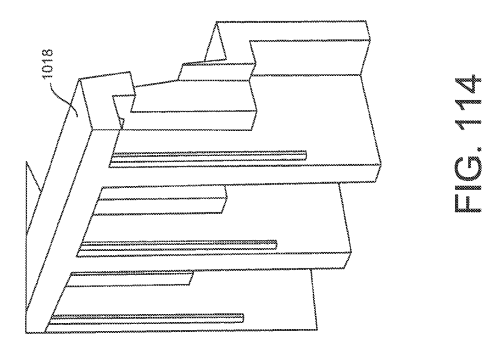


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Apr. 19, 2016



## TISSUE PENETRATION DEVICE

# RELATED APPLICATIONS

This application claims the benefit of U.S. Ser. No. 60/758, 5774, filed Jan. 12, 2006. This application is also a continuation-in-part of U.S. Ser. No. 11/318,334, filed Dec. 22, 2005 now abandoned, which is a divisional of U.S. Ser. No. 10/127, 395, filed Apr. 19, 2002 now U.S. Pat. No. 7,025,774. All of the above-referenced applications are fully incorporated 10 herein by reference.

#### BACKGROUND

#### 1. Technical Field

The technical field relates to analyte sampling devices, and more specifically, methods and devices for storing analyte sampling and measurement devices in a safe, usable condition.

### 2. Description of the Related Art

Lancing devices are known in the medical health-care products industry for piercing the skin to produce blood for analysis. Typically, a drop of blood for this type of analysis is obtained by making a small incision in the fingertip, creating a small wound, which generates a small blood droplet on the 25 surface of the skin.

Early methods of lancing included piercing or slicing the skin with a needle or razor. Current methods utilize lancing devices that contain a multitude of spring, cam and mass actuators to drive the lancet. These include cantilever springs, 30 diaphragms, coil springs, as well as gravity plumbs used to drive the lancet. The device may be held against the skin and mechanically triggered to ballistically launch the lancet. Unfortunately, the pain associated with each lancing event using known technology discourages patients from testing. In 35 addition to vibratory stimulation of the skin as the driver impacts the end of a launcher stop, known spring based devices have the possibility of firing lancets that harmonically oscillate against the patient tissue, causing multiple strikes due to recoil. This recoil and multiple strikes of the lancet is 40 one major impediment to patient compliance with a structured glucose monitoring regime.

Success rate generally encompasses the probability of producing a blood sample with one lancing action, which is sufficient in volume to perform the desired analytical test. The 45 blood may appear spontaneously at the surface of the skin, or may be "milked" from the wound. Milking generally involves pressing the side of the digit, or in proximity of the wound to express the blood to the surface. In traditional methods, the blood droplet produced by the lancing action must reach the 50 surface of the skin to be viable for testing.

When using existing methods, blood often flows from the cut blood vessels but is then trapped below the surface of the skin, forming a hematoma. In other instances, a wound is created, but no blood flows from the wound. In either case, the 55 lancing process cannot be combined with the sample acquisition and testing step. Spontaneous blood droplet generation with current mechanical launching system varies between launcher types but on average it is about 50% of lancet strikes, which would be spontaneous. Otherwise milking is required 60 to yield blood. Mechanical launchers are unlikely to provide the means for integrated sample acquisition and testing if one out of every two strikes does not yield a spontaneous blood sample.

Many diabetic patients (insulin dependent) are required to 65 self-test for blood glucose levels five to six times daily. The large number of steps required in traditional methods of glu-

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cose testing ranging from lancing, to milking of blood, applying blood to the test strip, and getting the measurements from the test strip discourages many diabetic patients from testing their blood glucose levels as often as recommended. Tight control of plasma glucose through frequent testing is therefore mandatory for disease management. The pain associated with each lancing event further discourages patients from testing. Additionally, the wound channel left on the patient by known systems may also be of a size that discourages those who are active with their hands or who are worried about healing of those wound channels from testing their glucose levels

Another problem frequently encountered by patients who must use lancing equipment to obtain and analyze blood samples is the amount of manual dexterity and hand-eye coordination required to properly operate the lancing and sample testing equipment due to retinopathies and neuropathies particularly, severe in elderly diabetic patients. For those patients, operating existing lancet and sample testing equipment can be a challenge. Once a blood droplet is created, that droplet must then be guided into a receiving channel of a small test strip or the like. If the sample placement on the strip is unsuccessful, repetition of the entire procedure including re-lancing the skin to obtain a new blood droplet is necessary.

Early methods of using test strips required a relatively substantial volume of blood to obtain an accurate glucose measurement. This large blood requirement made the monitoring experience a painful one for the user since the user may need to lance deeper than comfortable to obtain sufficient blood generation. Alternatively, if insufficient blood is spontaneously generated, the user may need to "milk" the wound to squeeze enough blood to the skin surface. Neither method is desirable as they take additional user effort and may be painful. The discomfort and inconvenience associated with such lancing events may deter a user from testing their blood glucose levels in a rigorous manner sufficient to control their diabetes.

A further impediment to patient compliance is the technique for storing these analyte sampling and analyte detecting devices. The devices used to measure analyte levels are typically stored in a humidity controlled or other safe environment to maintain the device shelf life. This often involves using a variety of containers, some for the test strips and some for the lancets. The introduction of multiple storage devices and the cumbersome design may discourage users from keeping their equipment in a usable condition, further degrading user test compliance and measurement accuracy.

There is a need for a device to measure analyte levels with improved humidity control. There is a further need for a device to measure analyte levels that includes desiccant that is external to penetrating members. There is a further need for a device, which can reliably, repeatedly and painlessly generate spontaneous blood samples. In addition, a method for performing analytical testing on a sample that does not require a high degree of manual dexterity or hand-eye coordination is required. Integrating sample generation (lancing) with sample testing (sample to test strip) will result in a simple one-step testing procedure resulting in better disease management through increased compliance with self testing regimes.

#### **SUMMARY**

Accordingly, an object of the present invention is to provide a disposable cartridge that has separate and integrated penetrating member and analyte sensor housings.

Another object of the present invention is to provide a disposable cartridge that has separation of sterilization of penetrating members from the analyte sensors.

A further object of the present invention is to provide a disposable cartridge with sterilization of penetrating members and maintenance of analyte sensors in a dry condition.

Yet another object of the present invention is to provide a disposable cartridge with alignment of penetrating members to analyte sensors enables that upon penetration by a penetrating member of a skin surface, blood flows into an analyte 10

Still another object of the present invention is to provide a disposable cartridge that measures analyte levels with improved humidity control.

Another object of the present invention is to provide a 15 of the tissue penetration device of FIG. 20. disposable cartridge to measure analyte levels that includes desiccant external to penetrating members.

A further object of the present invention is to provide a disposable cartridge that can reliably, repeatedly and painlessly generate spontaneous blood samples.

These and other objects of the present invention are achieved in a body fluid sampling system for use on a tissue site. A drive force generator is coupled to penetrating members. The penetrating members are at least partially housed in a penetrating member housing. The penetrating member 25 housing has openings for the penetrating members. The penetrating members are launched along a launch path by the drive force generator through an opening in the penetrating member housing. Analyte sensors are housed in a analyte sensor housing. Each analyte sensor is associated with a 30 penetrating member. The analyte sensor housing and the penetrating member housing are coupled to provide that each analyte sensor is positioned out of a launch path of a penetrating member. The analyte sensor housing includes desiccant pockets for receiving desiccant plugs.

### BRIEF DESCRIPTION OF THE DRAWINGS

FIGS. 1-3 are graphs of lancet velocity versus position for force drivers.

FIG. 4 illustrates an embodiment of a controllable force driver in the form of a flat electric lancet driver that has a solenoid-type configuration.

FIG. 5 illustrates an embodiment of a controllable force 45 driver in the form of a cylindrical electric lancet driver using a coiled solenoid-type configuration.

FIG. 6 illustrates a displacement over time profile of a lancet driven by a harmonic spring/mass system.

FIG. 7 illustrates the velocity over time profile of a lancet 50 ing from the skin of a patient's finger. driver by a harmonic spring/mass system.

FIG. 8 illustrates a displacement over time profile of an embodiment of a controllable force driver.

FIG. 9 illustrates a velocity over time profile of an embodiment of a controllable force driver.

FIG. 10 illustrates the lancet needle partially retracted, after severing blood vessels; blood is shown following the needle in the wound tract.

FIG. 11 illustrates blood following the lancet needle to the skin surface, maintaining an open wound tract.

FIG. 12 is a diagrammatic view illustrating a controlled feed-back loop.

FIG. 13 is a graph of force vs. time during the advancement and retraction of a lancet showing some characteristic phases of a lancing cycle.

FIG. 14 illustrates a lancet tip showing features, which can affect lancing pain, blood volume, and success rate.

FIG. 15 illustrates an embodiment of a lancet tip.

FIG. 16 is a graph showing displacement of a lancet over

FIG. 17 is a graph showing an embodiment of a velocity profile, which includes the velocity of a lancet over time including reduced velocity during retraction of the lancet.

FIG. 18 illustrates the tip of an embodiment of a lancet before, during and after the creation of an incision braced with a helix.

FIG. 19 illustrates a finger wound tract braced with an elastomer embodiment.

FIG. 20 is a perspective view of a tissue penetration device having features of the invention.

FIG. 21 is an elevation view in partial longitudinal section

FIG. 22 is an elevation view in partial section of an alternative embodiment.

FIG. 23 is a transverse cross sectional view of the tissue penetration device of FIG. 21 taken along lines 23-23 of FIG. 20 21.

FIG. 24 is a transverse cross sectional view of the tissue penetration device of FIG. 21 taken along lines 24-24 of FIG.

FIG. 25 is a transverse cross sectional view of the tissue penetration device of FIG. 21 taken along lines 25-25 of FIG.

FIG. 26 is a transverse cross sectional view of the tissue penetration device of FIG. 21 taken along lines 26-26 of FIG. 21.

FIG. 27 is a side view of the drive coupler of the tissue penetration device of FIG. 21.

FIG. 28 is a front view of the drive coupler of the tissue penetration device of FIG. 21 with the lancet not shown for purposes of illustration.

FIGS. 29A-29C show a flowchart illustrating a lancet control method.

FIG. 30 is a diagrammatic view of a patient's finger and a lancet tip moving toward the skin of the finger.

FIG. 31 is a diagrammatic view of a patient's finger and the embodiments of spring driven, cam driven, and controllable 40 lancet tip making contact with the skin of a patient's finger.

> FIG. 32 is a diagrammatic view of the lancet tip depressing the skin of a patient's finger.

> FIG. 33 is a diagrammatic view of the lancet tip further depressing the skin of a patient's finger.

> FIG. 34 is a diagrammatic view of the lancet tip penetrating the skin of a patient's finger.

> FIG. 35 is a diagrammatic view of the lancet tip penetrating the skin of a patient's finger to a desired depth.

> FIG. 36 is a diagrammatic view of the lancet tip withdraw-

FIGS. 37-41 illustrate a method of tissue penetration that may measure elastic recoil of the skin.

FIG. 42 is a graphical representation of position and velocity vs. time for a lancing cycle.

FIG. 43 illustrates a sectional view of the layers of skin with a lancet disposed therein.

FIG. 44 is a graphical representation of velocity vs. position of a lancing cycle.

FIG. 45 is a graphical representation of velocity vs. time of 60 a lancing cycle.

FIG. 46 is an elevation view in partial longitudinal section of an alternative embodiment of a driver coil pack and position analyte sensor.

FIG. 47 is a perspective view of a flat coil driver having 65 features of the invention.

FIG. 48 is an exploded view of the flat coil driver of FIG.

- FIG. 49 is an elevational view in partial longitudinal section of a tapered driver coil pack having features of the inven-
- FIG. 50 is a transverse cross sectional view of the tapered coil driver pack of FIG. 49 taken along lines 50-50 in FIG. 49.
- FIG. 51 shows an embodiment of a sampling module which houses a lancet and sample reservoir.
- FIG. 52 shows a housing that includes a driver and a chamber where the module shown in FIG. 51 can be loaded.
- FIG. 53 shows a tissue penetrating sampling device with the module loaded into the housing.
- FIG. 54 shows an alternate embodiment of a lancet con-
- FIG. 55 illustrates an embodiment of a sample input port, 15 sample reservoir and ergonomically contoured finger contact
- FIG. 56 illustrates the tissue penetration sampling device during a lancing event.
- FIG. 57 illustrates a thermal sample analyte sensor having 20 a sample detection element near a surface over which a fluid may flow and an alternative position for a sampled detection element that would be exposed to a fluid flowing across the surface.
- FIG. **58** shows a configuration of a thermal sample analyte 25 sensor with a sample detection element that includes a separate heating element.
- FIG. 59 depicts three thermal sample detectors such as that shown in FIG. 58 with sample detection elements located near each other alongside a surface.
- FIG. 60 illustrates thermal sample analyte sensors positioned relative to a channel having an analysis site.
- FIG. 61 shows thermal sample analyte sensors with sample detection analyzers positioned relative to analysis sites 35 arranged in an array on a surface.
- FIG. 62 schematically illustrates a sampling module device including several possible configurations of thermal sample analyte sensors including sample detection elements posiregions.
- FIG. 63 illustrates a tissue penetration sampling device having features of the invention.
- FIG. 64 is a top view in partial section of a sampling module of the tissue penetration sampling device of FIG. 63. 45
- FIG. 65 is a cross sectional view through line 65-65 of the sampling module shown in FIG. 64.
- FIG. 66 schematically depicts a sectional view of an alternative embodiment of the sampling module.
- FIG. 67 depicts a portion of the sampling module surround- 50 loaded in the lateral slot. ing a sampling port.
- FIGS. 68-70 show in sectional view one implementation of a spring powered lancet driver in three different positions during use of the lancet driver.
- FIG. 71 illustrates an embodiment of a tissue penetration 55 sampling device having features of the invention.
- FIG. 72 shows a top surface of a cartridge that includes multiple sampling modules.
- FIG. 73 shows in partial section a sampling module of the sampling cartridge positioned in a reader device.
- FIG. 74 is a perspective view in partial section of a tissue penetration sampling device with a cartridge of sampling modules.
- FIG. 75 is a front view in partial section of the tissue penetration sampling device of FIG. 56.
- FIG. 76 is a top view of the tissue penetration sampling device of FIG. 75.

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- FIG. 77 is a perspective view of a section of a sampling module belt having a plurality of sampling modules connected in series by a sheet of flexible polymer.
- FIG. 78 is a perspective view of a single sampling module of the sampling module belt of FIG. 59.
- FIG. 79 is a bottom view of a section of the flexible polymer sheet of the sampling module of FIG. 78 illustrating the flexible conductors and contact points deposited on the bottom surface of the flexible polymer sheet.
- FIG. 80 is a perspective view of the body portion of the sampling module of FIG. 77 without the flexible polymer cover sheet or lancet.
- FIG. 81 is an enlarged portion of the body portion of the sampling module of FIG. 80 illustrating the input port, sample flow channel, analytical region, lancet channel and lancet guides of the sampling module.
- FIG. 82 is an enlarged elevational view of a portion of an alternative embodiment of a sampling module having a plurality of small volume analytical regions.
- FIG. 83 is a perspective view of a body portion of a lancet module that can house and guide a lancet without sampling or analytical functions.
- FIG. 84 is an elevational view of a drive coupler having a T-slot configured to accept a drive head of a lancet.
- FIG. **85** is an elevational view of the drive coupler of FIG. 84 from the side and illustrating the guide ramps of the drive coupler.
- FIG. 86 is a perspective view of the drive coupler of FIG. 84 with a lancet being loaded into the T-slot of the drive coupler.
- FIG. 87 is a perspective view of the drive coupler of FIG. 86 with the drive head of the lancet completely loaded into the T-slot of the drive coupler.
- FIG. 88 is a perspective view of a sampling module belt disposed within the T-slot of the drive coupler with a drive head of a lancet of one of the sampling modules loaded within the T-slot of the drive coupler.
- FIG. 89 is a perspective view of a sampling module cartioned relative to sample flow channels and analytical 40 tridge with the sampling modules arranged in a ring configuration.
  - FIG. 90 is a perspective view of a sampling module cartridge with the plurality of sampling modules arranged in a block matrix with lancet drive heads configured to mate with a drive coupler having adhesive coupling.
  - FIG. 91 is a side view of an alternative embodiment of a drive coupler having a lateral slot configured to accept the L-shaped drive head of the lancet that is disposed within a lancet module and shown with the L-shaped drive head
  - FIG. 92 is an exploded view of the drive coupler, lancet with L-shaped drive head and lancet module of FIG. 91.
  - FIG. 93 is a perspective view of the front of a lancet cartridge coupled to the distal end of a controlled electromagnetic driver.
  - FIG. 94 is an elevational front view of the lancet cartridge
    - FIG. 95 is a top view of the lancet cartridge of FIG. 93.
  - FIG. **96** is a perspective view of the lancet cartridge of FIG. 60 93 with a portion of the cartridge body and lancet receptacle not shown for purposes of illustration of the internal mecha-
    - FIGS. 97-101 illustrate an embodiment of an agent injection device.
  - FIGS. 102-106 illustrate an embodiment of a cartridge for use in sampling having a sampling cartridge body and a lancet cartridge body.

FIGS. 107-110 illustrate embodiments of a disposable cartridge of the present invention with a analyte sensor housing and a penetrating member housing.

FIGS. **111-114** illustrate various embodiments of sensor housings that can be utilized with embodiments of the present 5 invention.

#### DETAILED DESCRIPTION

Variations in skin thickness including the stratum corneum and hydration of the epidermis can yield different results between different users with existing tissue penetration devices, such as lancing devices wherein the tissue penetrating element of the tissue penetration device is a lancet. Many current devices rely on adjustable mechanical stops or damp- 15 ing, to control the lancet's depth of penetration.

Displacement velocity profiles for both spring driven and cam driven tissue penetration devices are shown in FIGS. 1 and 2, respectively. Velocity is plotted against displacement X of the lancet. FIG. 1 represents a displacement/velocity pro- 20 file typical of spring driven devices. The lancet exit velocity increases until the lancet hits the surface of the skin 10. Because of the tensile characteristics of the skin, it will bend or deform until the lancet tip cuts the surface 20, the lancet will then penetrate the skin until it reaches a full stop 30. At 25 this point displacement is maximal and reaches a limit of penetration and the lancet stops. Mechanical stops absorb excess energy from the driver and transfer it to the lancet. The energy stored in the spring can cause recoil resulting in multiple piercing as seen by the coiled profile in FIG. 1. This 30 results in unnecessary pain from the additional tissue penetration as well as from transferring vibratory energy into the skin and exciting nerve endings. Retraction of the lancet then occurs and the lancet exits the skin 40 to return into the housing. Velocity cannot be controlled in any meaningful way 35 for this type of spring-powered driver.

FIG. 2 shows a displacement/velocity profile for a cam driven driver, which is similar to that of FIG. 1, but because the return path is specified in the cam configuration, there is no possibility of multiple tissue penetrations from one actuation. Cam based drivers can offer some level of control of lancet velocity vs. displacement, but not enough to achieve many desirable displacement/velocity profiles.

Advantages are achieved by utilizing a controllable force driver to drive a lancet, such as a driver, powered by electro-45 magnetic energy. A controllable driver can achieve a desired velocity versus position profile, such as that shown in FIG. 3. Embodiments of the present invention allow for the ability to accurately control depth of penetration, to control lancet penetration and withdrawal velocity, and therefore reduce the 50 pain perceived when cutting into the skin. Embodiments of the invention include a controllable driver that can be used with a feedback loop with a position analyte sensor to control the power delivered to the lancet, which can optimize the velocity and displacement profile to compensate for variations in skin thickness.

Pain reduction can be achieved by using a rapid lancet cutting speed, which is facilitated by the use of a lightweight lancet. The rapid cuffing minimizes the shock waves produced when the lancet strikes the skin in addition to compressing the skin for efficient cutting. If a controllable driver is used, the need for a mechanical stop can be eliminated. Due to the very light mass of the lancet and lack of a mechanical stop, there is little or no vibrational energy transferred to the finger during cutting.

The lancing devices such as those whose velocity versus position profiles are shown in FIGS. 1 and 2 typically yield

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50% spontaneous blood. In addition, some lancing events are unsuccessful and yield no blood, even on milking the finger. A spontaneous blood droplet generation is dependent on reaching the blood capillaries and venuoles, which yield the blood sample. It is therefore an issue of correct depth of penetration of the cutting device. Due to variations in skin thickness and hydration, some types of skin will deform more before cutting starts, and hence the actual depth of penetration will be less, resulting in less capillaries and venuoles cut. A controllable force driver can control the depth of penetration of a lancet and hence improve the spontaneity of blood yield. Furthermore, the use of a controllable force driver can allow for slow retraction of the lancet (slower than the cutting velocity) resulting in improved success rate due to the would channel remaining open for the free passage of blood to the surface of the skin.

Spontaneous blood yield occurs when blood from the cut vessels flow up the wound tract to the surface of the skin, where it can be collected and tested. Tissue elasticity parameters may force the wound tract to close behind the retracting lancet preventing the blood from reaching the surface. If however, the lancet were to be withdrawn slowly from the wound tract, thus keeping the wound open, blood could flow up the patent channel behind the tip of the lancet as it is being withdrawn (ref. FIGS. 10 and 11). Hence the ability to control the lancet speed into and out of the wound allows the device to compensate for changes in skin thickness and variations in skin hydration and thereby achieves spontaneous blood yield with maximum success rate while minimizing pain.

An electromagnetic driver can be coupled directly to the lancet minimizing the mass of the lancet and allowing the driver to bring the lancet to a stop at a predetermined depth without the use of a mechanical stop. Alternatively, if a mechanical stop is required for positive positioning, the energy transferred to the stop can be minimized. The electromagnetic driver allows programmable control over the velocity vs. position profile of the entire lancing process including timing the start of the lancet, tracking the lancet position, measuring the lancet velocity, controlling the distal stop acceleration, and controlling the skin penetration depth.

Referring to FIG. 4, an embodiment of a tissue penetration device is shown. The tissue penetration device includes a controllable force driver in the form of an electromagnetic driver, which can be used to drive a lancet. The term Lancet, as used herein, generally includes any sharp or blunt member, preferably having a relatively low mass, used to puncture the skin for the purpose of cutting blood vessels and allowing blood to flow to the surface of the skin. The term Electromagnetic driver, as used herein, generally includes any device that moves or drives a tissue penetrating element, such as a lancet under an electrically or magnetically induced force. FIG. 4 is a partially exploded view of an embodiment of an electromagnetic driver. The top half of the driver is shown assembled. The bottom half of the driver is shown exploded for illustrative purposes.

FIG. 4 shows the inner insulating housing 22 separated from the stationary housing or PC board 20, and the lancet 24 and flag 26 assembly separated from the inner insulating housing 22 for illustrative purposes. In addition, only four rivets 18 are shown as attached to the inner insulating housing 22 and separated from the PC board 20. In an embodiment, each coil drive field core in the PC board located in the PC Board 20 and 30 is connected to the inner insulating housing 22 and 32 with rivets.

The electromagnetic driver has a moving part comprising a lancet assembly with a lancet 24 and a magnetically permeable flag 26 attached at the proximal or drive end and a

stationary part comprising a stationary housing assembly with electric field coils arranged so that they produce a balanced field at the flag to reduce or eliminate any net lateral force on the flag. The electric field coils are generally one or more metal coils, which generate a magnetic field when electric current passes through the coil. The iron flag is a flat or enlarged piece of magnetic material, which increases the surface area of the lancet assembly to enhance the magnetic forces generated between the proximal end of the lancet and a magnetic field produced by the field coils. The combined mass of the lancet and the iron flag can be minimized to facilitate rapid acceleration for introduction into the skin of a patient, to reduce the impact when the lancet stops in the skin, and to facilitate prompt velocity profile changes throughout the sampling cycle.

The stationary housing assembly consists of a PC board 20, a lower inner insulating housing 22, an upper inner insulating housing 32, an upper PC board 30, and rivets 18 assembled into a single unit. The lower and upper inner insulating housing 22 and 32 are relieved to form a slot so that lancet assembly can be slid into the driver assembly from the side perpendicular to the direction of the lancet's advancement and retraction. This allows the disposal of the lancet assembly and reuse of the stationary housing assembly with another lancet assembly while avoiding accidental lancet launches during replacement.

The electric field coils in the upper and lower stationary housing 20 and 30 are fabricated in a multi-layer printed circuit (PC) board. They may also be conventionally wound 30 wire coils. A Teflon® material, or other low friction insulating material is used to construct the lower and upper inner insulating housing 22 and 32. Each insulating housing is mounted on the PC board to provide electrical insulation and physical protection, as well as to provide a low-friction guide for the 35 lancet. The lower and upper inner insulating housing 22 and 32 provide a reference surface with a small gap so that the lancet assembly 24 and 26 can align with the drive field coils in the PC board for good magnetic coupling.

Rivets 18 connect the lower inner insulating housing 22 to the lower stationary housing 20 and are made of magnetically permeable material such as ferrite or steel, which serves to concentrate the magnetic field. This mirrors the construction of the upper inner insulating housing 32 and upper stationary housing 30. These rivets form the poles of the electric field 45 coils. The PC board is fabricated with multiple layers of coils or with multiple boards. Each layer supports spiral traces around a central hole. Alternate layers spiral from the center outwards or from the edges inward. In this way each layer connects via simple feed-through holes, and the current 50 always travels in the same direction, summing the ampereturns.

The PC boards within the lower and upper stationary housings 20 and 30 are connected to the lower and upper inner insulating housings 22 and 32 with the rivets 18. The lower 35 and upper inner insulating housings 22 and 32 expose the rivet heads on opposite ends of the slot where the lancet assembly 24 and 26 travels. The magnetic field lines from each rivet create magnetic poles at the rivet heads. An iron bar on the opposite side of the PC board within each of the lower and upper stationary housing 20 and 30 completes the magnetic circuit by connecting the rivets. Any fastener made of magnetically permeable material such as iron or steel can be used in place of the rivets. A single component made of magnetically permeable material and formed in a horseshoe shape can 65 be used in place of the rivet/screw and iron bar assembly. In operation, the magnetically permeable flag 26 attached to the

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lancet 24 is divided into slits and bars 34. The slit patterns are staggered so that coils can drive the flag 26 in two, three or more phases.

Both lower and upper PC boards 20 and 30 contain drive coils so that there is a symmetrical magnetic field above and below the flag 26. When the pair of PC boards is turned on, a magnetic field is established around the bars between the slits of the magnetically permeable iron on the flag 26. The bars of the flag experience a force that tends to move the magnetically permeable material to a position minimizing the number and length of magnetic field lines and conducting the magnetic field lines between the magnetic poles.

When a bar of the flag 26 is centered between the rivets 18 of a magnetic pole, there is no net force on the flag, and any disturbing force is resisted by imbalance in the field. This embodiment of the device operates on a principle similar to that of a solenoid. Solenoids cannot push by repelling iron; they can only pull by attracting the iron into a minimum energy position. The slits 34 on one side of the flag 26 are offset with respect to the other side by approximately one half of the pitch of the poles. By alternately activating the coils on each side of the PC board, the lancet assembly can be moved with respect to the stationary housing assembly. The direction of travel is established by selectively energizing the coils adjacent the metal flag on the lancet assembly. Alternatively, a three phase, three-pole design or a shading coil that is offset by one-quarter pitch establishes the direction of travel. The lower and upper PC boards 20 and 30 shown in FIG. 4 contain electric field coils, which drive the lancet assembly and the circuitry for controlling the entire electromagnetic driver.

The embodiment described above generally uses the principles of a magnetic attraction drive, similar to commonly available circular stepper motors (Hurst Manufacturing BA Series motor, or "Electrical Engineering Handbook" Second edition p 1472-1474, 1997). These references are hereby incorporated by reference. Other embodiments can include a linear induction drive that uses a changing magnetic field to induce electric currents in the lancet assembly. These induced currents produce a secondary magnetic field that repels the primary field and applies a net force on the lancet assembly. The linear induction drive uses an electrical drive control that sweeps a magnetic field from pole to pole, propelling the lancet before it. Varying the rate of the sweep and the magnitude of the field by altering the driving voltage and frequency controls the force applied to the lancet assembly and its velocity.

The arrangement of the coils and rivets to concentrate the magnetic flux also applies to the induction design creating a growing magnetic field as the electric current in the field switches on. This growing magnetic field creates an opposing electric current in the conductive flag. In a linear induction motor the flag is electrically conductive, and its magnetic properties are unimportant. Copper or aluminum are materials that can be used for the conductive flags. Copper is generally used because of its good electrical conductivity. The opposing electrical field produces an opposing magnetic field that repels the field of the coils. By phasing the power of the coils, a moving field can be generated which pushes the flag along just below the synchronous speed of the coils. By controlling the rate of sweep, and by generating multiple sweeps, the flag can be moved at a desired speed.

FIG. 5 shows another embodiment of a solenoid type electromagnetic driver that is capable of driving an iron core or slug mounted to the lancet assembly using a direct current (DC) power supply. The electromagnetic driver includes a driver coil pack that is divided into three separate coils along the path of the lancet, two end coils and a middle coil. Direct

current is alternated to the coils to advance and retract the lancet. Although the driver coil pack is shown with three coils, any suitable number of coils may be used, for example, 4, 5, 6, 7 or more coils may be used.

The stationary iron housing **40** contains the driver coil pack 5 with a first coil **52** is flanked by iron spacers **50** which concentrate the magnetic flux at the inner diameter creating magnetic poles. The inner insulating housing **48** isolates the lancet **42** and iron core **46** from the coils and provides a smooth, low friction guide surface. The lancet guide **44** further centers the 10 lancet **42** and iron core **46**. The lancet **42** is protracted and retracted by alternating the current between the first coil **52**, the middle coil, and the third coil to attract the iron core **46**. Reversing the coil sequence and attracting the core and lancet back into the housing retracts the lancet. The lancet guide **44** 15 also serves as a stop for the iron core **46** mounted to the lancet **42** 

As discussed above, tissue penetration devices which employ spring or cam driving methods have a symmetrical or nearly symmetrical actuation displacement and velocity profiles on the advancement and retraction of the lancet as shown in FIGS. 6 and 7. In most of the available lancet devices, once the launch is initiated, the stored energy determines the velocity profile until the energy is dissipated. Controlling impact, retraction velocity, and dwell time of the lancet within the 25 tissue can be useful in order to achieve a high success rate while accommodating variations in skin properties and minimize pain. Advantages can be achieved by taking into account that tissue dwell time is related to the amount of skin deformation as the lancet tries to puncture the surface of the skin 30 and variance in skin deformation from patient to patient based on skin hydration.

The ability to control velocity and depth of penetration can be achieved by use of a controllable force driver where feedback is an integral part of driver control. Such drivers can 35 control either metal or polymeric lancets or any other type of tissue penetration element. The dynamic control of such a driver is illustrated in FIG. 8 which illustrates an embodiment of a controlled displacement profile and FIG. 9 which illustrates an embodiment of a the controlled velocity profile. 40 These are compared to FIGS. 6 and 7, which illustrate embodiments of displacement and velocity profiles, respectively, of a harmonic spring/mass powered driver.

Reduced pain can be achieved by using impact velocities of greater than 2 m/s entry of a tissue penetrating element, such 45 as a lancet, into tissue.

Retraction of the lancet at a low velocity following the sectioning of the venuole/capillary mesh allows the blood to flood the wound tract and flow freely to the surface, thus using the lancet to keep the channel open during retraction as shown in FIGS. 10 and 11. Low-velocity retraction of the lancet near the wound flap prevents the wound flap from sealing off the channel. Thus, the ability to slow the lancet retraction directly contributes to increasing the success rate of obtaining blood. Increasing the sampling success rate to near 100% can be 55 important to the combination of sampling and acquisition into an integrated sampling module such as an integrated glucose-sampling module, which incorporates a glucose test strip.

Referring again to FIG. 5, the lancet and lancet driver are configured so that feedback control is based on lancet displacement, velocity, or acceleration. The feedback control information relating to the actual lancet path is returned to a processor such as that illustrated in FIG. 12 that regulates the energy to the driver, thereby precisely controlling the lancet throughout its advancement and retraction. The driver may be 65 driven by electric current, which includes direct current and alternating current.

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In FIG. 5, the electromagnetic driver shown is capable of driving an iron core or slug mounted to the lancet assembly using a direct current (DC) power supply and is also capable of determining the position of the iron core by measuring magnetic coupling between the core and the coils. The coils can be used in pairs to draw the iron core into the driver coil pack. As one of the coils is switched on, the corresponding induced current in the adjacent coil can be monitored. The strength of this induced current is related to the degree of magnetic coupling provided by the iron core, and can be used to infer the position of the core and hence, the relative position of the lancet.

After a period of time, the drive voltage can be turned off, allowing the coils to relax, and then the cycle is repeated. The degree of magnetic coupling between the coils is converted electronically to a proportional DC voltage that is supplied to an analog-to-digital converter. The digitized position signal is then processed and compared to a desired "nominal" position by a central processing unit (CPU). The CPU to set the level and/or length of the next power pulse to the solenoid coils uses error between the actual and nominal positions.

In another embodiment, the driver coil pack has three coils consisting of a central driving coil flanked by balanced detection coils built into the driver assembly so that they surround an actuation or magnetically active region with the region centered on the middle coil at mid-stroke. When a current pulse is applied to the central coil, voltages are induced in the adjacent sense coils. If the sense coils are connected together so that their induced voltages oppose each other, the resulting signal will be positive for deflection from mid-stroke in one direction, negative in the other direction, and zero at mid-stroke. This measuring technique is commonly used in Linear Variable Differential Transformers (LVDT). Lancet position is determined by measuring the electrical balance between the two sensing coils.

In another embodiment, a feedback loop can use a commercially available LED/photo transducer module such as the OPB703 manufactured by Optek Technology, Inc., 1215 W. Crosby Road, Carrollton, Tex., 75006 to determine the distance from the fixed module on the stationary housing to a reflective surface or target mounted on the lancet assembly. The LED acts as a light emitter to send light beams to the reflective surface, which in turn reflects the light back to the photo transducer, which acts as a light analyte sensor. Distances over the range of 4 mm or so are determined by measuring the intensity of the reflected light by the photo transducer. In another embodiment, a feedback loop can use a magnetically permeable region on the lancet shaft itself as the core of a Linear Variable Differential Transformer (LVDT).

A permeable region created by selectively annealing a portion of the lancet shaft, or by including a component in the lancet assembly, such as ferrite, with sufficient magnetic permeability to allow coupling between adjacent sensing coils. Coil size, number of windings, drive current, signal amplification, and air gap to the permeable region are specified in the design process. In another embodiment, the feedback control supplies a piezoelectric driver, superimposing a high frequency oscillation on the basic displacement profile. The piezoelectric driver provides improved cutting efficiency and reduces pain by allowing the lancet to "saw" its way into the tissue or to destroy cells with cavitation energy generated by the high frequency of vibration of the advancing edge of the lancet. The drive power to the piezoelectric driver is monitored for an impedance shift as the device interacts with the target tissue. The resulting force measurement, coupled with the known mass of the lancet is used to determine lancet acceleration, velocity, and position.

FIG. 12 illustrates the operation of a feedback loop using a processor. The processor 60 stores profiles 62 in non-volatile memory. A user inputs information 64 about the desired circumstances or parameters for a lancing event. The processor 60 selects a driver profile 62 from a set of alternative driver 5 profiles that have been preprogrammed in the processor 60 based on typical or desired tissue penetration device performance determined through testing at the factory or as programmed in by the operator. The processor 60 may customize by either scaling or modifying the profile based on additional user input information 64. Once the processor has chosen and customized the profile, the processor 60 is ready to modulate the power from the power supply 66 to the lancet driver 68 through an amplifier 70. The processor 60 measures the location of the lancet 72 using a position sensing mechanism 74 15 through an analog to digital converter 76. Examples of position sensing mechanisms have been described in the embodiments above. The processor 60 calculates the movement of the lancet by comparing the actual profile of the lancet to the predetermined profile. The processor 60 modulates the power 20 to the lancet driver 68 through a signal generator 78, which controls the amplifier 70 so that the actual profile of the lancet does not exceed the predetermined profile by more than a preset error limit. The error limit is the accuracy in the control of the lancet.

After the lancing event, the processor 60 can allow the user to rank the results of the lancing event. The processor 60 stores these results and constructs a database 80 for the individual user. Using the database 80, the processor 60 calculates the profile traits such as degree of painlessness, success 30 rate, and blood volume for various profiles 62 depending on user input information 64 to optimize the profile to the individual user for subsequent lancing cycles. These profile traits depend on the characteristic phases of lancet advancement and retraction. The processor 60 uses these calculations to 35 optimize profiles 62 for each user. In addition to user input information 64, an internal clock allows storage in the database 80 of information such as the time of day to generate a time stamp for the lancing event and the time between lancing events to anticipate the user's diurnal needs. The database 40 stores information and statistics for each user and each profile that particular user uses.

In addition to varying the profiles, the processor **60** can be used to calculate the appropriate lancet diameter and geometry necessary to realize the blood volume required by the 45 user. For example, if the user requires a 1-5 micro liter volume of blood, the processor selects a 200 micron diameter lancet to achieve these results. For each class of lancet, both diameter and lancet tip geometry, is stored in the processor to correspond with upper and lower limits of attainable blood volume based on the predetermined displacement and velocity profiles.

The lancing device is capable of prompting the user for information at the beginning and the end of the lancing event to more adequately suit the user. The goal is to either change 55 to a different profile or modify an existing profile. Once the profile is set, the force driving the lancet is varied during advancement and retraction to follow the profile. The method of lancing using the lancing device comprises selecting a profile, lancing according to the selected profile, determining 60 lancing profile traits for each characteristic phase of the lancing cycle, and optimizing profile traits for subsequent lancing events.

FIG. 13 shows an embodiment of the characteristic phases of lancet advancement and retraction on a graph of force 65 versus time illustrating the force exerted by the lancet driver on the lancet to achieve the desired displacement and velocity

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profile. The characteristic phases are the lancet introduction phase A-C where the lancet is longitudinally advanced into the skin, the lancet rest phase D where the lancet terminates its longitudinal movement reaching its maximum depth and becoming relatively stationary, and the lancet retraction phase E-G where the lancet is longitudinally retracted out of the skin. The duration of the lancet retraction phase E-G is longer than the duration of the lancet introduction phase A-C, which in turn is longer than the duration of the lancet rest phase D.

The introduction phase further comprises a lancet launch phase prior to A when the lancet is longitudinally moving through air toward the skin, a tissue contact phase at the beginning of A when the distal end of the lancet makes initial contact with the skin, a tissue deformation phase A when the skin bends depending on its elastic properties which are related to hydration and thickness, a tissue lancing phase which comprises when the lancet hits the inflection point on the skin and begins to cut the skin B and the lancet continues cutting the skin C. The lancet rest phase D is the limit of the penetration of the lancet into the skin. Pain is reduced by minimizing the duration of the lancet introduction phase A-C so that there is a fast incision to a certain penetration depth regardless of the duration of the deformation phase A and inflection point cutting B which will vary from user to user. Success rate is increased by measuring the exact depth of penetration from inflection point B to the limit of penetration in the lancet rest phase D. This measurement allows the lancet to always, or at least reliably, hit the capillary beds which are a known distance underneath the surface of the skin.

The lancet retraction phase further comprises a primary retraction phase E when the skin pushes the lancet out of the wound tract, a secondary retraction phase F when the lancet starts to become dislodged and pulls in the opposite direction of the skin, and lancet exit phase G when the lancet becomes free of the skin. Primary retraction is the result of exerting a decreasing force to pull the lancet out of the skin as the lancet pulls away from the finger. Secondary retraction is the result of exerting a force in the opposite direction to dislodge the lancet. Control is necessary to keep the wound tract open as blood flows up the wound tract. Blood volume is increased by using a uniform velocity to retract the lancet during the lancet retraction phase E-G regardless of the force required for the primary retraction phase E or secondary retraction phase F, either of which may vary from user to user depending on the properties of the user's skin.

FIG. 14 shows a standard industry lancet for glucose testing which has a three-facet geometry. Taking a rod of diameter 114 and grinding 8 degrees to the plane of the primary axis to create the primary facet 110 produces the lancet 116. The secondary facets 112 are then created by rotating the shaft of the needle 15 degrees, and then rolling over 12 degrees to the plane of the primary facet. Other possible geometry's require altering the lancet's production parameters such as shaft diameter, angles, and translation distance.

FIG. 15 illustrates facet and tip geometry 120 and 122, diameter 124, and depth 126 which are significant factors in reducing pain, blood volume and success rate. It is known that additional cutting by the lancet is achieved by increasing the shear percentage or ratio of the primary to secondary facets, which when combined with reducing the lancet's diameter reduces skin tear and penetration force and gives the perception of less pain. Overall success rate of blood yield, however, also depends on a variety of factors, including the existence of facets, facet geometry, and skin anatomy.

FIG. 16 shows another embodiment of displacement versus time profile of a lancet for a controlled lancet retraction.

FIG. 17 shows the velocity vs. time profile of the lancet for the controlled retraction of FIG. 16. The lancet driver controls lancet displacement and velocity at several steps in the lancing cycle, including when the lancet cuts the blood vessels to allow blood to pool 130, and as the lancet retracts, regulating 5 the retraction rate to allow the blood to flood the wound tract while keeping the wound flap from sealing the channel 132 to permit blood to exit the wound.

In addition to slow retraction of a tissue-penetrating element in order to hold the wound open to allow blood to escape to the skin surface, other methods are contemplated. FIG. 18 shows the use of an embodiment of the invention, which includes a retractable coil on the lancet tip. A coiled helix or tube 140 is attached externally to lancet 116 with the freedom to slide such that when the lancet penetrates the skin 150, the 15 helix or tube 140 follows the trajectory of the lancet 116. The helix begins the lancing cycle coiled around the facets and shaft of the lancet 144. As the lancet penetrates the skin, the helix braces the wound tract around the lancet 146. As the lancet retracts, the helix remains to brace open the wound 20 tract, keeping the wound tract from collapsing and keeping the surface skin flap from closing 148. This allows blood 152 to pool and flow up the channel to the surface of the skin. The helix is then retracted as the lancet pulls the helix to the point where the helix is decompressed to the point where the diam- 25 eter of the helix becomes less than the diameter of the wound tract and becomes dislodged from the skin.

The tube or helix **140** is made of wire or metal of the type commonly used in angioplasty stents such as stainless steel, nickel titanium alloy or the like. Alternatively the tube or 30 helix **140** or a ring can be made of a biodegradable material, which braces the wound tract by becoming lodged in the skin. Biodegradation is completed within seconds or minutes of insertion, allowing adequate time for blood to pool and flow up the wound tract. Biodegradation is activated by heat, moisture, or pH from the skin.

Alternatively, the wound could be held open by coating the lancet with a powder or other granular substance. The powder coats the wound tract and keeps it open when the lancet is withdrawn. The powder or other granular substance can be a 40 coarse bed of microspheres or capsules which hold the channel open while allowing blood to flow through the porous interstices.

In another embodiment the wound can be held open using a two-part needle, the outer part in the shape of a "U" and the 45 inner part filling the "U." After creating the wound the inner needle is withdrawn leaving an open channel, rather like the plugs that are commonly used for withdrawing sap from maple trees.

FIG. 19 shows a further embodiment of a method and device for facilitating blood flow utilizing an elastomer to coat the wound. This method uses an elastomer 154, such as silicon rubber, to coat or brace the wound tract 156 by covering and stretching the surface of the finger 158. The elastomer 154 is applied to the finger 158 prior to lancing. After a short delay, the lancet (not shown) then penetrates the elastomer 154 and the skin on the surface of the finger 158 as is seen in 160. Blood is allowed to pool and rise to the surface while the elastomer 154 braces the wound tract 156 as is seen in 162 and 164. Other known mechanisms for increasing the success rate of blood yield after lancing can include creating a vacuum, suctioning the wound, applying an adhesive strip, vibration while cutting, or initiating a second lance if the first is unsuccessful.

FIG. 20 illustrates an embodiment of a tissue penetration 65 device, more specifically, a lancing device 180 that includes a controllable driver 179 coupled to a tissue penetration ele-

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ment. The lancing device 180 has a proximal end 181 and a distal end 182. At the distal end 182 is the tissue penetration element in the form of a lancet 183, which is coupled to an elongate coupler shaft 184 by a drive coupler 185. The elongate coupler shaft 184 has a proximal end 186 and a distal end 187. A driver coil pack 188 is disposed about the elongate coupler shaft 184 proximal of the lancet 183. A position analyte sensor 191 is disposed about a proximal portion 192 of the elongate coupler shaft 184 and an electrical conductor 194 electrically couples a processor 193 to the position analyte sensor 191. The elongate coupler shaft 184 driven by the driver coil pack 188 controlled by the position analyte sensor 191 and processor 193 form the controllable driver, specifically, a controllable electromagnetic driver.

Referring to FIG. 21, the lancing device 180 can be seen in more detail, in partial longitudinal section. The lancet 183 has a proximal end 195 and a distal end 196 with a sharpened point at the distal end 196 of the lancet 183 and a drive head 198 disposed at the proximal end 195 of the lancet 183. A lancet shaft 201 is disposed between the drive head 198 and the sharpened point 197. The lancet shaft 201 may be comprised of stainless steel, or any other suitable material or alloy and have a transverse dimension of about 0.1 to about 0.4 mm. The lancet shaft may have a length of about 3 mm to about 50 mm, specifically, about 15 mm to about 20 mm. The drive head 198 of the lancet 183 is an enlarged portion having a transverse dimension greater than a transverse dimension of the lancet shaft 201 distal of the drive head 198. This configuration allows the drive head 198 to be mechanically captured by the drive coupler 185. The drive head 198 may have a transverse dimension of about 0.5 to about 2 mm.

A magnetic member 202 is secured to the elongate coupler shaft 184 proximal of the drive coupler 185 on a distal portion 203 of the elongate coupler shaft 184. The magnetic member 202 is a substantially cylindrical piece of magnetic material having an axial lumen 204 extending the length of the magnetic member 202. The magnetic member 202 has an outer transverse dimension that allows the magnetic member 202 to slide easily within an axial lumen 205 of a low friction, possibly lubricious, polymer guide tube 205' disposed within the driver coil pack 188. The magnetic member 202 may have an outer transverse dimension of about 1.0 to about 5.0 mm, specifically, about 2.3 to about 2.5 mm. The magnetic member 202 may have a length of about 3.0 to about 5.0 mm, specifically, about 4.7 to about 4.9 mm. The magnetic member 202 can be made from a variety of magnetic materials including ferrous metals such as ferrous steel, iron, ferrite, or the like. The magnetic member 202 may be secured to the distal portion 203 of the elongate coupler shaft 184 by a variety of methods including adhesive or epoxy bonding, welding, crimping or any other suitable method.

Proximal of the magnetic member 202, an optical encoder flag 206 is secured to the elongate coupler shaft 184. The optical encoder flag 206 is configured to move within a slot 207 in the position analyte sensor 191. The slot 207 of the position analyte sensor 191 is formed between a first body portion 208 and a second body portion 209 of the position analyte sensor 191. The slot 207 may have separation width of about 1.5 to about 2.0 mm. The optical encoder flag 206 can have a length of about 14 to about 18 mm, a width of about 3 to about 5 mm and a thickness of about 0.04 to about 0.06 mm.

The optical encoder flag 206 interacts with various optical beams generated by LEDs disposed on or in the position analyte sensor body portions 208 and 209 in a predetermined manner. The interaction of the optical beams generated by the LEDs of the position analyte sensor 191 generates a signal that indicates the longitudinal position of the optical flag 206

relative to the position analyte sensor 191 with a substantially high degree of resolution. The resolution of the position analyte sensor 191 may be about 200 to about 400 cycles per inch, specifically, about 350 to about 370 cycles per inch. The position analyte sensor 191 may have a speed response time (position/time resolution) of 0 to about 120,000 Hz, where one dark and light stripe of the flag constitutes one Hertz, or cycle per second. The position of the optical encoder flag 206 relative to the magnetic member 202, driver coil pack 188 and position analyte sensor 191 is such that the optical encoder 191 can provide precise positional information about the lancet 183 over the entire length of the lancet's power stroke.

An optical encoder that is suitable for the position analyte sensor 191 is a linear optical incremental encoder, model HEDS 9200, manufactured by Agilent Technologies. The 15 model HEDS 9200 may have a length of about 20 to about 30 mm, a width of about 8 to about 12 mm, and a height of about 9 to about 11 mm. Although the position analyte sensor 191 illustrated is a linear optical incremental encoder, other suitable position analyte sensor embodiments could be used, 20 provided they posses the requisite positional resolution and time response. The HEDS 9200 is a two channel device where the channels are 90 degrees out of phase with each other. This results in a resolution of four times the basic cycle of the flag. These quadrature outputs make it possible for the processor to 25 determine the direction of lancet travel. Other suitable position analyte sensors include capacitive encoders, analog reflective analyte sensors, such as the reflective position analyte sensor discussed above, and the like.

A coupler shaft guide 211 is disposed towards the proximal 30 end 181 of the lancing device 180. The guide 211 has a guide lumen 212 disposed in the guide 211 to slidingly accept the proximal portion 192 of the elongate coupler shaft 184. The guide 211 keeps the elongate coupler shaft 184 centered horizontally and vertically in the slot 202 of the optical 35 encoder 191.

The driver coil pack 188, position analyte sensor 191 and coupler shaft guide 211 are all secured to a base 213. The base 213 is longitudinally coextensive with the driver coil pack 188, position analyte sensor 191 and coupler shaft guide 211. 40 The base 213 can take the form of a rectangular piece of metal or polymer, or may be a more elaborate housing with recesses, which are configured to accept the various components of the lancing device 180.

As discussed above, the magnetic member 202 is config- 45 ured to slide within an axial lumen 205 of the driver coil pack **188**. The driver coil pack **188** includes a most distal first coil 214, a second coil 215, which is axially disposed between the first coil 214 and a third coil 216, and a proximal-most fourth coil 217. Each of the first coil 214, second coil 215, third coil 50 216 and fourth coil 217 has an axial lumen. The axial lumens of the first through fourth coils are configured to be coaxial with the axial lumens of the other coils and together form the axial lumen 205 of the driver coil pack 188 as a whole. Axially adjacent each of the coils 214-217 is a magnetic disk or 55 washer 218 that augments completion of the magnetic circuit of the coils 214-217 during a lancing cycle of the device 180. The magnetic washers 218 of the embodiment of FIG. 21 are made of ferrous steel but could be made of any other suitable magnetic material, such as iron or ferrite. The outer shell 189 of the driver coil pack 188 is also made of iron or steel to complete the magnetic path around the coils and between the washers 218. The magnetic washers 218 have an outer diameter commensurate with an outer diameter of the driver coil pack 188 of about 4.0 to about 8.0 mm. The magnetic washers 218 have an axial thickness of about 0.05, to about 0.4 mm, specifically, about 0.15 to about 0.25 mm.

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Wrapping or winding an elongate electrical conductor 221 about an axial lumen until a sufficient number of windings have been achieved forms the coils 214-217. The elongate electrical conductor 221 is generally an insulated solid copper wire with a small outer transverse dimension of about 0.06 mm to about 0.88 mm, specifically, about 0.3 mm to about 0.5 mm. In one embodiment, 32 gauge copper wire is used for the coils 214-217. The number of windings for each of the coils 214-217 of the driver pack 188 may vary with the size of the coil, but for some embodiments each coil 214-217 may have about 30 to about 80 turns, specifically, about 50 to about 60 turns. Each coil 214-217 can have an axial length of about 1.0 to about 3.0 mm, specifically, about 1.8 to about 2.0 mm. Each coil 214-217 can have an outer transverse dimension or diameter of about 4.0, to about 2.0 mm, specifically, about 9.0 to about 12.0 mm. The axial lumen 205 can have a transverse dimension of about 1.0 to about 3.0 mm.

It may be advantageous in some driver coil 188 embodiments to replace one or more of the coils with permanent magnets, which produce a magnetic field similar to that of the coils when the coils are activated. In particular, it may be desirable in some embodiments to replace the second coil 215, the third coil 216 or both with permanent magnets. In addition, it may be advantageous to position a permanent magnet at or near the proximal end of the coil driver pack in order to provide fixed magnet zeroing function for the magnetic member (Adams magnetic Products 23A0002 flexible magnet material (800) 747-7543).

FIGS. 20 and 21 show a permanent bar magnet 219 disposed on the proximal end of the driver coil pack 188. As shown in FIG. 21, the bar magnet 219 is arranged so as to have one end disposed adjacent the travel path of the magnetic member 202 and has a polarity configured so as to attract the magnetic member 202 in a centered position with respect to the bar magnet 219. Note that the polymer guide tube 205' can be configured to extend proximally to insulate the inward radial surface of the bar magnet 219 from an outer surface of the magnetic member 202. This arrangement allows the magnetic member 219 and thus the elongate coupler shaft 184 to be attracted to and held in a zero point or rest position without the consumption of electrical energy from the power supply 225

Having a fixed zero or start point for the elongate coupler shaft 184 and lancet 183 can be critical to properly controlling the depth of penetration of the lancet 183 as well as other lancing parameters. This can be because some methods of depth penetration control for a controllable driver measure the acceleration and displacement of the elongate coupler shaft 184 and lancet 183 from a known start position. If the distance of the lancet tip 196 from the target tissue is known, acceleration and displacement of the lancet is known and the start position of the lancet is know, the time and position of tissue contact and depth of penetration can be determined by the processor 193.

Any number of configurations for a magnetic bar 219 can be used for the purposes discussed above. In particular, a second permanent bar magnet (not shown) could be added to the proximal end of the driver coil pack 188 with the magnetic fields of the two bar magnets configured to complement each other. In addition, a disc magnet 219' could be used as illustrated in FIG. 22. Disc magnet 219' is shown disposed at the proximal end of the driver coiled pack 188 with a polymer non-magnetic disc 219" disposed between the proximal-most coil 217 and disc magnet 219' and positions disc magnet 219' away from the proximal end of the proximal-most coil 217. The polymer non-magnetic disc spacer 219" is used so that the magnetic member 202 can be centered in a zero or start

position slightly proximal of the proximal-most coil 217 of the driver coil pack 188. This allows the magnetic member to be attracted by the proximal-most coil 217 at the initiation of the lancing cycle instead of being passive in the forward drive portion of the lancing cycle.

An inner lumen of the polymer non-magnetic disc 219" can be configured to allow the magnetic member 202 to pass axially there through while an inner lumen of the disc magnet 219' can be configured to allow the elongate coupler shaft 184 to pass through but not large enough for the magnetic member 202 to pass through. This results in the magnetic member 202 being attracted to the disc magnet 219' and coming to rest with the proximal surface of the magnetic member 202 against a distal surface of the disc magnet 219'. This arrangement provides for a positive and repeatable stop for the magnetic member, and hence the lancet. A similar configuration could also be used for the bar magnet 219 discussed above.

Typically, when the electrical current in the coils 214-217 of the driver coil pack 188 is off, a magnetic member 202 20 made of soft iron is attracted to the bar magnet 219 or disc magnet 219'. The magnetic field of the driver coil pack 188 and the bar magnet 219 or disc magnet 219', or any other suitable magnet, can be configured such that when the electrical current in the coils 214-217 is turned on, the leakage 25 magnetic field from the coils 214-217 has the same polarity as the bar magnet 219 or disc magnet 219'. This results in a magnetic force that repels the magnetic member 202 from the bar magnet 219 or disc magnet 219' and attracts the magnetic member 202 to the activated coils 214-217. For this configuration, the bar magnet 219 or disc magnet thus act to facilitate acceleration of the magnetic member 202 as opposed to working against the acceleration.

Electrical conductors 222 couple the driver coil pack 188 with the processor 193 which can be configured or programmed to control the current flow in the coils 214-217 of the driver coil pack 188 based on position feedback from the position analyte sensor 191, which is coupled to the processor 193 by electrical conductors 194. A power source 225 is electrically coupled to the processor 193 and provides electrical power to operate the processor 193 and power the coil driver pack 188. The power source 225 may be one or more batteries that provide direct current power to the 193 processor.

FIG. 23 shows a transverse cross sectional view of drive 45 coupler 185 in more detail. The drive head 198 of the lancet 183 is disposed within the drive coupler 185 with a first retaining rail 226 and second retaining rail 227 capturing the drive head 198 while allowing the drive head 198 to be inserted laterally into the drive coupler 185 and retracted 50 laterally with minimal mechanical resistance. The drive coupler 185 may optionally be configured to include snap ridges 228 which allow the drive head 198 to be laterally inserted and retracted, but keep the drive head 198 from falling out of the drive coupler 185 unless a predetermined amount of exter- 55 nally applied lateral force is applied to the drive head 198 of the lancet 183 towards the lateral opening 231 of the drive coupler 185. FIG. 27 shows an enlarged side view into the coupler opening 231 of the drive coupler 185 showing the snap ridges 228 disposed in the lateral opening 231 and the 60 retaining rails 226 and 227. FIG. 28 shows an enlarged front view of the drive coupler 185. The drive coupler 185 can be made from an alloy such as stainless steel, titanium or aluminum, but may also be made from a suitable polymer such as ABS, PVC, polycarbonate plastic or the like. The drive coupler may be open on both sides allowing the drive head and lancet to pass through.

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Referring to FIG. 24, the magnetic member 202 is disposed about and secured to the elongate coupler shaft 184. The magnetic member 202 is disposed within the axial lumen 232 of the fourth coil 217. The driver coil pack 188 is secured to the base 213. In FIG. 25 the position analyte sensor 191 is secured to the base 213 with the first body portion 208 of the position analyte sensor 191 disposed opposite the second body portion 209 of the position analyte sensor 191 with the first and second body portions 208 and 209 of the position analyte sensor 191 separated by the gap or slot 207. The elongate coupler shaft 184 is slidably disposed within the gap 207 between the first and second body portions 208 and 209 of the position analyte sensor 191. The optical encoder flag 206 is secured to the elongate coupler shaft 184 and disposed between the first body portion 208 and second body portion 209 of the position analyte sensor 191. Referring to FIG. 26, the proximal portion 192 of the elongate coupler shaft 184 is disposed within the guide lumen 212 of the coupler shaft guide 211. The guide lumen 212 of the coupler shaft guide 211 may be lined with a low friction material such as Teflon® or the like to reduce friction of the elongate coupler shaft 184 during the power stroke of the lancing device 180.

Referring to FIGS. 29A-29C, a flow diagram is shown that describes the operations performed by the processor 193 in controlling the lancet 183 of the lancing device 180 discussed above during an operating cycle. FIGS. 30-36 illustrate the interaction of the lancet 183 and skin 233 of the patient's finger 234 during an operation cycle of the lancet device 183. The processor 193 operates under control of programming steps that are stored in an associated memory. When the programming steps are executed, the processor 193 performs operations as described herein. Thus, the programming steps implement the functionality of the operations described with respect to the flow diagram of FIG. 29. The processor 193 can receive the programming steps from a program product stored in recordable media, including a direct access program product storage device such as a hard drive or flash ROM, a removable program product storage device such as a floppy disk, or in any other manner known to those of skill in the art. The processor 193 can also download the programming steps through a network connection or serial connection.

In the first operation, represented by the flow diagram box numbered 245 in FIG. 29A, the processor 193 initializes values that it stores in memory relating to control of the lancet, such as variables that it uses to keep track of the controllable driver 179 during movement. For example, the processor may set a clock value to zero and a lancet position value to zero or to some other initial value. The processor 193 may also cause power to be removed from the coil pack 188 for a period of time, such as for about 10 ms, to allow any residual flux to dissipate from the coils.

In the initialization operation, the processor 193 also causes the lancet to assume an initial stationary position. When in the initial stationary position, the lancet 183 is typically fully retracted such that the magnetic member 202 is positioned substantially adjacent the fourth coil 217 of the driver coil pack 188, shown in FIG. 21 above. The processor 193 can move the lancet 183 to the initial stationary position by pulsing an electrical current to the fourth coil 217 to thereby attract the magnetic member 202 on the lancet 183 to the fourth coil 217. Alternatively, the magnetic member can be positioned in the initial stationary position by virtue of a permanent magnet, such as bar magnet 219, disc magnet 219 or any other suitable magnet as discussed above with regard to the tissue penetration device illustrated in FIGS. 20 and 21.

In the next operation, represented by the flow diagram box numbered 247, the processor 193 energizes one or more of the

coils in the coil pack 188. This should cause the lancet 183 to begin to move (i.e., achieve a non-zero speed) toward the skin target 233. The processor 193 then determines whether or not the lancet is indeed moving, as represented by the decision box numbered 249. The processor 193 can determine whether the lancet 183 is moving by monitoring the position of the lancet 183 to determine whether the position changes over time. The processor 193 can monitor the position of the lancet 183 by keeping track of the position of the optical encoder

If the processor 193 determines (via timeout without motion events) that the lancet 183 is not moving (a "No" result from the decision box 249), then the process proceeds 15 to the operation represented by the flow diagram box numbered 253, where the processor deems that an error condition is present. This means that some error in the system is causing the lancet 183 not to move. The error may be mechanical, electrical, or software related. For example, the lancet 183 20 may be stuck in the stationary position because something is impeding its movement.

flag 206 secured to the elongate coupler shaft 184 wherein the

encoder 191 produces a signal coupled to the processor 193

that indicates the spatial position of the lancet 183.

If the processor 193 determines that the lancet 183 is indeed moving (a "Yes" result from the decision box numbered **249**), then the process proceeds to the operation repre- 25 sented by the flow diagram box numbered 257. In this operation, the processor 193 causes the lancet 183 to continue to accelerate and launch toward the skin target 233, as indicated by the arrow 235 in FIG. 30. The processor 193 can achieve acceleration of the lancet 183 by sending an electrical current 30 to an appropriate coil 214-217 such that the coil 214-217 exerts an attractive magnetic launching force on the magnetic member 202 and causes the magnetic member 202 and the lancet 183 coupled thereto to move in a desired direction. For example, the processor 193 can cause an electrical current to 35 be sent to the third coil 216 so that the third coil 216 attracts the magnetic member 202 and causes the magnetic member 202 to move from a position adjacent the fourth coil 217 toward the third coil 216. The processor preferably determines which coil 214-217 should be used to attract the mag- 40 netic member 202 based on the position of the magnetic member 202 relative to the coils 214-217. In this manner, the processor 193 provides a controlled force to the lancet that controls the movement of the lancet.

During this operation, the processor 193 periodically or continually monitors the position and/or velocity of the lancet 183. In keeping track of the velocity and position of the lancet 183 as the lancet 183 moves towards the patient's skin 233 or other tissue, the processor 193 also monitors and adjusts the electrical current to the coils 214-217. In some embodiments, 50 the processor 193 applies current to an appropriate coil 214-217 such that the lancet 183 continues to move according to a desired direction and acceleration. In the instant case, the processor 193 applies current to the appropriate coil 214-217 that will cause the lancet 183 to continue to move in the direction of the patient's skin 233 or other tissue to be penetrated

The processor 193 may successively transition the current between coils 214-217 so that as the magnetic member 202 moves past a particular coil 214-217, the processor 193 then 60 shuts off current to that coil 214-217 and then applies current to another coil 214-217 that will attract the magnetic member 202 and cause the magnetic member 202 to continue to move in the desired direction. In transitioning current between the coils 214-217, the processor 193 can take into account various 65 factors, including the speed of the lancet 183, the position of the lancet 183 relative to the coils 214-217, the number of

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coils 214-217, and the level of current to be applied to the coils 214-217 to achieve a desired speed or acceleration.

In the next operation, the processor 193 determines whether the cutting or distal end tip 196 of the lancet 183 has contacted the patient's skin 233, as shown in FIG. 31 and as represented by the decision box numbered 265 in FIG. 29B. The processor 193 may determine whether the lancet 183 has made contact with the target tissue 233 by a variety of methods, including some that rely on parameters which are measured prior to initiation of a lancing cycle and other methods that are adaptable to use during a lancing cycle without any predetermined parameters.

In one embodiment, the processor 193 determines that the skin has been contacted when the end tip 196 of the lancet 183 has moved a predetermined distance with respect to its initial position. If the distance from the tip 961 of the lancet 183 to the target tissue 233 is known prior to initiation of lancet 183 movement, the initial position of the lancet 183 is fixed and known, and the movement and position of the lancet 183 can be accurately measured during a lancing cycle, then the position and time of lancet contact can be determined.

This method requires an accurate measurement of the distance between the lancet tip 196 and the patient's skin 233 when the lancet 183 is in the zero time or initial position. This can be accomplished in a number of ways. One way is to control all of the mechanical parameters that influence the distance from the lancet tip 196 to the patient's tissue or a surface of the lancing device 180 that will contact the patient's skin 233. This could include the start position of the magnetic member 202, magnetic path tolerance, magnetic member 202 dimensions, driver coil pack 188 location within the lancing device 180 as a whole, length of the elongate coupling shaft 184, placement of the magnetic member 202 on the elongate coupling shaft 184, length of the lancet 183 etc.

If all these parameters, as well as others can be suitably controlled in manufacturing with a tolerance stack-up that is acceptable, then the distance from the lancet tip 196 to the target tissue 233 can be determined at the time of manufacture of the lancing device 180. The distance could then be programmed into the memory of the processor 193. If an adjustable feature is added to the lancing device 180, such as an adjustable length elongate coupling shaft 184, this can accommodate variations in all of the parameters noted above, except length of the lancet 183. An electronic alternative to this mechanical approach would be to calibrate a stored memory contact point into the memory of the processor 193 during manufacture based on the mechanical parameters described above.

In another embodiment, moving the lancet tip 196 to the target tissue 233 very slowly and gently touching the skin 233 prior to actuation can accomplish the distance from the lancet tip 196 to the tissue 233. The position analyte sensor can accurately measure the distance from the initialization point to the point of contact, where the resistance to advancement of the lancet 183 stops the lancet movement. The lancet 183 is then retracted to the initialization point having measured the distance to the target tissue 233 without creating any discomfort to the user.

In another embodiment, the processor 193 may use software to determine whether the lancet 183 has made contact with the patients skin 233 by measuring for a sudden reduction in velocity of the lancet 183 due to friction or resistance imposed on the lancet 183 by the patient's skin 233. The optical encoder 191 measures displacement of the lancet 183. The position output data provides input to the interrupt input of the processor 193. The processor 193 also has a timer

capable of measuring the time between interrupts. The distance between interrupts is known for the optical encoder 191, so the velocity of the lancet 183 can be calculated by dividing the distance between interrupts by the time between the interrupts.

This method requires that velocity losses to the lancet 183 and elongate coupler 184 assembly due to friction are known to an acceptable level so that these velocity losses and resulting deceleration can be accounted for when establishing a deceleration threshold above which contact between lancet 10 tip 196 and target tissue 233 will be presumed. This same concept can be implemented in many ways. For example, rather than monitoring the velocity of the lancet 183, if the processor 193 is controlling the lancet driver in order to maintain a fixed velocity, the power to the driver 188 could be 15 monitored. If an amount of power above a predetermined threshold is required in order to maintain a constant velocity, then contact between the tip of the lancet 196 and the skin 233 could be presumed.

In yet another embodiment, the processor 193 determines 20 skin 233 contact by the lancet 183 by detection of an acoustic signal produced by the tip 196 of the lancet 183 as it strikes the patient's skin 233. Detection of the acoustic signal can be measured by an acoustic detector 236 placed in contact with the patient's skin 233 adjacent a lancet penetration site 237, as 25 shown in FIG. 31. Suitable acoustic detectors 236 include piezo electric transducers, microphones and the like. The acoustic detector 236 transmits an electrical signal generated by the acoustic signal to the processor 193 via electrical conductors 238. In another embodiment, contact of the lancet 30 183 with the patient's skin 233 can be determined by measurement of electrical continuity in a circuit that includes the lancet 183, the patient's finger 234 and an electrical contact pad 240 that is disposed on the patient's skin 233 adjacent the contact site 237 of the lancet 183, as shown in FIG. 31. In this 35 embodiment, as soon as the lancet 183 contacts the patients skin 233, the circuit 239 is completed and current flows through the circuit 239. Completion of the circuit 239 can then be detected by the processor 193 to confirm skin 233 contact by the lancet 183.

If the lancet 183 has not contacted the target skin 233, then the process proceeds to a timeout operation, as represented by the decision box numbered 267 in FIG. 29B. In the timeout operation, the processor 193 waits a predetermined time period. If the timeout period has not yet elapsed (a "No" 45 outcome from the decision box 267), then the processor continues to monitor whether the lancet has contacted the target skin 233. The processor 193 preferably continues to monitor the position and speed of the lancet 183, as well as the electrical current to the appropriate coil 214-217 to maintain the 50 desired lancet 183 movement.

If the timeout period elapses without the lancet 183 contacting the skin (a "Yes" output from the decision box 267), then it is deemed that the lancet 183 will not contact the skin and the process proceeds to a withdraw phase, where the 55 lancet is withdrawn away from the skin 233, as discussed more fully below. The lancet 183 may not have contacted the target skin 233 for a variety of reasons, such as if the patient removed the skin 233 from the lancing device or if something obstructed the lancet 183 prior to it contacting the skin.

The processor 193 may also proceed to the withdraw phase prior to skin contact for other reasons. For example, at some point after initiation of movement of the lancet 183, the processor 193 may determine that the forward acceleration of the lancet 183 towards the patient's skin 233 should be stopped or 65 that current to all coils 214-217 should be shut down. This can occur, for example, if it is determined that the lancet 183 has

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achieved sufficient forward velocity, but has not yet contacted the skin 233. In one embodiment, the average penetration velocity of the lancet 183 from the point of contact with the skin to the point of maximum penetration may be about 2.0 to about 10.0 m/s, specifically, about 3.8 to about 4.2 m/s. In another embodiment, the average penetration velocity of the lancet may be from about 2 to about 8 meters per second, specifically, about 2 to about 4 m/s.

The processor 193 can also proceed to the withdraw phase if it is determined that the lancet 183 has fully extended to the end of the power stroke of the operation cycle of lancing procedure. In other words, the process may proceed to withdraw phase when an axial center 241 of the magnetic member 202 has moved distal of an axial center 242 of the first coil 214 as show in FIG. 21. In this situation, any continued power to any of the coils 214-217 of the driver coil pack 188 serves to decelerate the magnetic member 202 and thus the lancet 183. In this regard, the processor 193 considers the length of the lancet 183 (which can be stored in memory) the position of the lancet 183 relative to the magnetic member 202, as well as the distance that the lancet 183 has traveled.

With reference again to the decision box 265 in FIG. 29B, if the processor 193 determines that the lancet 183 has contacted the skin 233 (a "Yes" outcome from the decision box 265), then the processor 193 can adjust the speed of the lancet 183 or the power delivered to the lancet 183 for skin penetration to overcome any frictional forces on the lancet 183 in order to maintain a desired penetration velocity of the lancet. The flow diagram box numbered 267 represents this.

As the velocity of the lancet 183 is maintained after contact with the skin 233, the distal tip 196 of the lancet 183 will first begin to depress or tent the contacted skin 237 and the skin 233 adjacent the lancet 183 to form a tented portion 243 as shown in FIG. 32 and further shown in FIG. 33. As the lancet 183 continues to move in lancet 183 will eventually begin to penetrate the skin 233, as shown in FIG. 34. Once penetration of the skin 233 begins, the static force at the distal tip 196 of the lancet 183 from the skin 233 will become a dynamic cutting force, which is generally less than the static tip force. As a result in the reduction of force on the distal tip 196 of the lancet 183 upon initiation of cutting, the tented portion 243 of the skin 233 adjacent the distal tip 196 of the lancet 183 which had been depressed as shown in FIGS. 32 and 24 will spring back as shown in FIG. 34.

In the next operation, represented by the decision box numbered 271 in FIG. 29B, the processor 193 determines whether the distal end 196 of the lancet 183 has reached a brake depth. The brake depth is the skin penetration depth for which the processor 193 determines that deceleration of the lancet 183 is to be initiated in order to achieve a desired final penetration depth 244 of the lancet 183 as show in FIG. 35. The brake depth may be pre-determined and programmed into the processor's memory, or the processor 193 may dynamically determine the brake depth during the actuation. The amount of penetration of the lancet 183 in the skin 233 of the patient may be measured during the operation cycle of the lancet device 180. In addition, as discussed above, the penetration depth necessary for successfully obtaining a useable sample can depend on the amount of tenting of the skin 233 60 during the lancing cycle. The amount of tenting of the patient's skin 233 can in turn depend on the tissue characteristics of the patient such as elasticity, hydration etc. A method for determining these characteristics is discussed below with regard to skin 233 tenting measurements during the lancing cycle and illustrated in FIGS. 37-41.

Penetration measurement can be carried out by a variety of methods that are not dependent on measurement of tenting of

specifically, about 50 to about 200 milliseconds. For other embodiments, the settling time may be about 1 to about 200 milliseconds.

It is at this stage of the lancing cycle that a software method

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the patient's skin. In one embodiment, the penetration depth of the lancet 183 in the patient's skin 233 is measured by monitoring the amount of capacitance between the lancet 183 and the patients skin 233. In this embodiment, a circuit includes the lancet 183, the patient's finger 234, the processor 5 193 and electrical conductors connecting these elements. As the lancet 183 penetrates the patient's skin 233, the greater the amount of penetration, the greater the surface contact area between the lancet 183 and the patient's skin 233. As the contact area increases, so does the capacitance between the skin 233 and the lancet 183. The increased capacitance can be easily measured by the processor 193 using methods known in the art and penetration depth can then be correlated to the amount of capacitance. The same method can be used by measuring the electrical resistance between the lancet 183 15 and the patient's skin.

It is at this stage of the lancing cycle that a software method can be used to measure the amount of tenting of the patient's skin 233 and thus determine the skin 233 characteristics such as elasticity, hydration and others. Referring to FIGS. 37-41, a lancet 183 is illustrated in various phases of a lancing cycle with target tissue 233. FIG. 37 shows tip 196 of lancet 183 making initial contact with the skin 233 at the point of initial impact.

If the brake depth has not yet been reached, then a "No" results from the decision box 271 and the process proceeds to the timeout operation represented by the flow diagram box numbered 273. In the timeout operation, the processor 193 waits a predetermined time period. If the timeout period has not yet elapsed (a "No" outcome from the decision box 273), then the processor continues to monitor whether the brake depth has been reached. If the timeout period elapses without the lancet 183 achieving the brake depth (a "Yes" output from 25 the decision box 273), then the processor 193 deems that the lancet 183 will not reach the brake depth and the process proceeds to the withdraw phase, which is discussed more fully below. This may occur, for example, if the lancet 183 is stuck at a certain depth.

FIG. 38 illustrates an enlarged view of the lancet 183 making initial contact with the tissue 233 shown in FIG. 37. In FIG. 39, the lancet tip 196 has depressed or tented the skin 233 prior to penetration over a distance of X, as indicated by the arrow labeled X in FIG. 39. In FIG. 40, the lancet 183 has reached the full length of the cutting power stroke and is at maximum displacement. In this position, the lancet tip 196 has penetrated the tissue 233 a distance of Y, as indicated by the arrow labeled Y in FIG. 39. As can be seen from comparing FIG. 38 with FIG. 40, the lancet tip 196 was displaced a total distance of X plus Y from the time initial contact with the skin 233 was made to the time the lancet tip 196 reached its maximum extension as shown in FIG. 40. However, the lancet tip 196 has only penetrated the skin 233 a distance Y because of the tenting phenomenon.

With reference again to the decision box numbered 271 in FIG. 29B, if the lancet does reach the brake depth (a "Yes" result), then the process proceeds to the operation represented by the flow diagram box numbered 275. In this operation, the processor 193 causes a braking force to be applied to the 35 lancet to thereby reduce the speed of the lancet 183 to achieve a desired amount of final skin penetration depth 244, as shown in FIG. 26. Note that FIGS. 32 and 33 illustrate the lancet making contact with the patient's skin and deforming or depressing the skin prior to any substantial penetration of the 40 skin. The speed of the lancet 183 is preferably reduced to a value below a desired threshold and is ultimately reduced to zero. The processor 193 can reduce the speed of the lancet 183 by causing a current to be sent to a 214-217 coil that will exert an attractive braking force on the magnetic member 202 in a 45 proximal direction away from the patients tissue or skin 233, as indicated by the arrow 290 in FIG. 36. Such a negative force reduces the forward or distally oriented speed of the lancet 183. The processor 193 can determine which coil 214-217 to energize based upon the position of the magnetic 50 member 202 with respect to the coils 214-217 of the driver coil pack 188, as indicated by the position analyte sensor 191.

At the end of the power stroke of the lancet 183, as discussed above with regard to FIG. 26 and box 279 of FIG. 29C, the processor 193 allows the lancet to settle for about 8 msec. It is during this settling time that the skin 233 rebounds or relaxes back to approximately its original configuration prior to contact by the lancet 183 as shown in FIG. 41. The lancet tip 196 is still buried in the skin to a depth of Y, as shown in FIG. 41, however the elastic recoil of the tissue has displaced the lancet rearward or retrograde to the point of inelastic tenting that is indicated by the arrows Z in FIG. 41. During the rearward displacement of the lancet 183 due to the elastic tenting of the tissue 233, the processor reads and stores the position data generated by the position analyte sensor 191 and thus measures the amount of elastic tenting, which is the difference between X and Z.

In the next operation, the process proceeds to the withdraw phase, as represented by the flow diagram box numbered 277. The withdraw phase begins with the operation represented by the flow diagram box numbered 279 in FIG. 29C. Here, the processor 193 allows the lancet 183 to settle at a position of maximum skin penetration 244, as shown in FIG. 35. In this regard, the processor 193 waits until any motion in the lancet 183 (due to vibration from impact and spring energy stored in the skin, etc.) has stopped by monitoring changes in position of the lancet 183. The processor 193 preferably waits until several milliseconds (ms), such as on the order of about 8 ms, have passed with no changes in position of the lancet 183. This is an indication that movement of the lancet 183 has ceased entirely. In some embodiments, the lancet may be allowed to settle for about 1 to about 2000 milliseconds,

The tenting process and retrograde motion of the lancet **183** during the lancing cycle is illustrated graphically in FIG. 42 which shows both a velocity versus time graph and a position versus time graph of a lancet tip 196 during a lancing cycle that includes elastic and inelastic tenting. In FIG. 42, from point 0 to point A, the lancet 183 is being accelerated from the initialization position or zero position. From point A to point B, the lancet is in ballistic or coasting mode, with no additional power being delivered. At point B, the lancet tip 196 contacts the tissue 233 and begins to tent the skin 233 until it reaches a displacement C. As the lancet tip 196 approaches maximum displacement, braking force is applied to the lancet 183 until the lancet comes to a stop at point D. The lancet 183 then recoils in a retrograde direction during the settling phase of the lancing cycle indicated between D and E. Note that the magnitude of inelastic tenting indicated in FIG. 42 is exaggerated for purposes of illustration.

The amount of inelastic tenting indicated by Z tends to be fairly consistent and small compared to the magnitude of the elastic tenting. Generally, the amount of inelastic tenting Z can be about 120 to about 140 microns. As the magnitude of the inelastic tenting has a fairly constant value and is small compared to the magnitude of the elastic tenting for most patients and skin types, the value for the total amount of tenting for the penetration stroke of the lancet 183 is effectively equal to the rearward displacement of the lancet during

the settling phase as measured by the processor 193 plus a predetermined value for the inelastic recoil, such as 130 microns. Inelastic recoil for some embodiments can be about 100 to about 200 microns. The ability to measure the magnitude of skin 233 tenting for a patient is important to controlling the depth of penetration of the lancet tip 196 as the skin is generally known to vary in elasticity and other parameters due to age, time of day, level of hydration, gender and pathological state.

This value for total tenting for the lancing cycle can then be 10 used to determine the various characteristics of the patient's skin 233. Once a body of tenting data is obtained for a given patient, this data can be analyzed in order to predict the total lancet displacement, from the point of skin contact, necessary for a successful lancing procedure. This enables the tissue 15 penetration device to achieve a high success rate and minimize pain for the user. A rolling average table can be used to collect and store the tenting data for a patient with a pointer to the last entry in the table. When a new entry is input, it can replace the entry at the pointer and the pointer advances to the 20 next value. When an average is desired, all the values are added and the sum divided by the total number of entries by the processor 193. Similar techniques involving exponential decay (multiply by 0.95, add 0.05 times current value, etc.) are also possible.

With regard to tenting of skin 233 generally, some typical values relating to penetration depth are now discussed. FIG. 43 shows a cross sectional view of the layers of the skin 233. In order to reliably obtain a useable sample of blood from the skin 233, it is desirable to have the lancet tip 196 reach the 30 venuolar plexus of the skin. The stratum corneum is typically about 0.1 to about 0.6 mm thick and the distance from the top of the dermis to the venuole plexus can be from about 0.3 to about 1.4 mm. Elastic tenting can have a magnitude of up to about 2 mm or so, specifically, about 0.2 to about 2.0 mm, 35 with an average magnitude of about 1 mm. This means that the amount of lancet displacement necessary to overcome the tenting can have a magnitude greater than the thickness of skin necessary to penetrate in order to reach the venuolar plexus. The total lancet displacement from point of initial 40 skin contact may have an average value of about 1.7 to about 2.1 mm. In some embodiments, penetration depth and maximum penetration depth may be about 0.5 mm to about 5 mm, specifically, about 1 mm to about 3 mm. In some embodiments, a maximum penetration depth of about 0.5 to about 3 45 mm is useful.

Referring back to FIG. 29C, in the next operation, represented by the flow diagram box numbered 280 in FIG. 29C, the processor 193 causes a withdraw force to be exerted on the lancet 183 to retract the lancet 183 from the skin 233, as 50 shown by arrow 290 in FIG. 36. The processor 193 sends a current to an appropriate coil 214-217 so that the coil 214-217 exerts an attractive distally oriented force on the magnetic member 202, which should cause the lancet 183 to move backward in the desired direction. In some embodiments, the 55 lancet 183 is withdrawn with less force and a lower speed than the force and speed during the penetration portion of the operation cycle. Withdrawal speed of the lancet in some embodiments can be about 0.004 to about 0.5 m/s, specifically, about 0.006 to about 0.01 m/s. In other embodiments, useful withdrawal velocities can be about 0.001 to about 0.02 meters per second, specifically, about 0.001 to about 0.01 meters per second. For embodiments that use a relatively slow withdrawal velocity compared to the penetration velocity, the withdrawal velocity may up to about 0.02 meters per second. For such embodiments, a ratio of the average penetration velocity relative to the average withdrawal velocity can be

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about 100 to about 1000. In embodiments where a relatively slow withdrawal velocity is not important, a withdrawal velocity of about 2 to about 10 meters per second may be used.

In the next operation, the processor 193 determines whether the lancet 183 is moving in the desired backward direction as a result of the force applied, as represented by the decision box numbered 281. If the processor 193 determines that the lancet 183 is not moving (a "No" result from the decision box 281), then the processor 193 continues to cause a force to be exerted on the lancet 183, as represented by the flow diagram box numbered 282. The processor 193 may cause a stronger force to be exerted on the lancet 183 or may just continue to apply the same amount of force. The processor then again determines whether the lancet is moving, as represented by the decision box numbered 283. If movement is still not detected (a "No" result from the decision box numbered 283), the processor 193 determines that an error condition is present, as represented by the flow diagram box numbered **284**. In such a situation, the processor preferably de-energizes the coils to remove force from the lancet, as the lack of movement may be an indication that the lancet is stuck in the skin of the patient and, therefore, that it may be undesirable to continue to attempt pull the lancet out of the skin.

With reference again to the decision boxes numbered 281 and 283 in FIG. 29C, if the processor 193 determines that the lancet is indeed moving in the desired backward direction away from the skin 233, then the process proceeds to the operation represented by the flow diagram box numbered 285. In this operation, the backward movement of the lancet 183 continues until the lancet distal end has been completely withdrawn from the patients skin 233. As discussed above, in some embodiments the lancet 183 is withdrawn with less force and a lower speed than the force and sped during the penetration portion of the operation cycle. The relatively slow withdrawal of the lancet 183 may allow the blood from the capillaries of the patient accessed by the lancet 183 to follow the lancet 183 during withdrawal and reach the skin surface to reliably produce a usable blood sample. The process then

Controlling the lancet motion over the operating cycle of the lancet **183** as discussed above allows a wide variety of lancet velocity profiles to be generated by the lancing device **180**. In particular, any of the lancet velocity profiles discussed above with regard to other embodiments can be achieved with the processor **193**, position analyte sensor **191** and driver coil pack **188** of the lancing device **180**.

Another example of an embodiment of a velocity profile for a lancet can be seen in FIGS. **44** and **45**, which illustrates a lancet profile with a fast entry velocity and a slow withdrawal velocity. FIG. **44** illustrates an embodiment of a lancing profile showing velocity of the lancet versus position. The lancing profile starts at zero time and position and shows acceleration of the lancet towards the tissue from the electromagnetic force generated from the electromagnetic driver. At point A, the power is shut off and the lancet **183** begins to coast until it reaches the skin **233** indicated by B at which point, the velocity begins to decrease. At point C, the lancet **183** has reached maximum displacement and settles momentarily, typically for a time of about 8 milliseconds.

A retrograde withdrawal force is then imposed on the lancet by the controllable driver, which is controlled by the processor to maintain a withdrawal velocity of no more than about 0.006 to about 0.01 meters/second. The same cycle is illustrated in the velocity versus time plot of FIG. 45 where the lancet is accelerated from the start point to point A. The lancet 183 coasts from A to B where the lancet tip 196 con-

tacts tissue 233. The lancet tip 196 then penetrates the tissue and slows with braking force eventually applied as the maximum penetration depth is approached. The lancet is stopped and settling between C and D. At D, the withdrawal phase begins and the lancet 183 is slowly withdrawn until it returns to the initialization point shown by E in FIG. 45. Note that retrograde recoil from elastic and inelastic tenting was not shown in the lancing profiles of FIGS. 44 and 45 for purpose of illustration and clarity.

In another embodiment, the withdrawal phase may use a dual speed profile, with the slow 0.006 to 0.01 meter per second speed used until the lancet is withdrawn past the contact point with the tissue, then a faster speed of 0.01 to 1 meters per second may be used to shorten the complete cycle.

Referring to FIG. 46, another embodiment of a lancing device including a controllable driver 294 with a driver coil pack 295, position analyte sensor and lancet 183 are shown. The lancet 297 has a proximal end 298 and a distal end 299 with a sharpened point at the distal end 299 of the lancet 297.

A magnetic member 301 disposed about and secured to a proximal end portion 302 of the lancet 297 with a lancet shaft 303 being disposed between the magnetic member 301 and the sharpened point 299. The lancet shaft 303 may be comprised of stainless steel, or any other suitable material or alloy.

The lancet shaft 303 may have a length of about 3 mm to about 50 mm specifically, about 5 mm to about 15 mm.

The magnetic member 301 is configured to slide within an axial lumen 304 of the driver coil pack 295. The driver coil pack 295 includes a most distal first coil 305, a second coil 30 306, which is axially disposed between the first coil 305 and a third coil 307, and a proximal-most fourth coil 308. Each of the first coil 305, second coil 306, third coil 307 and fourth coil 308 has an axial lumen. The axial lumens of the first through fourth coils 305-308 are configured to be coaxial 35 with the axial lumens of the other coils and together form the axial lumen 309 of the driver coil pack 295 as a whole. Axially adjacent each of the coils 305-308 is a magnetic disk or washer 310 that augments completion of the magnetic circuit of the coils 305-308 during a lancing cycle of the driven coil 40 pack 295. The magnetic washers 310 of the embodiment of FIG. 46 are made of ferrous steel but could be made of any other suitable magnetic material, such as iron or ferrite. The magnetic washers 310 have an outer diameter commensurate with an outer diameter of the driver coil pack 295 of about 4.0 45 to about 8.0 mm. The magnetic washers 310 have an axial thickness of about 0.05, to about 0.4 mm, specifically, about 0.15 to about 0.25 mm. The outer shell 294 of the coil pack is also made of iron or steel to complete the magnetic path around the coils and between the washers 310.

Wrapping or winding an elongate electrical conductor 311 about the axial lumen 309 until a sufficient number of windings have been achieved forms the coils 305-308. The elongate electrical conductor 311 is generally an insulated solid copper wire. The particular materials, dimensions number of 55 coil windings etc. of the coils 305-308, washers 310 and other components of the driver coil pack 295 can be the same or similar to the materials, dimensions number of coil windings etc. of the driver coil pack 188 discussed above.

Electrical conductors 312 couple the driver coil pack 295 60 with a processor 313 which can be configured or programmed to control the current flow in the coils 305-308 of the driver coil pack 295 based on position feedback from the position analyte sensor 296, which is coupled to the processor 313 by electrical conductors 315. A power source 316 is electrically 65 coupled to the processor 313 and provides electrical power to operate the processor 313 and power the driver coil pack 295.

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The power source 316 may be one or more batteries (not shown) that provide direct current power to the processor 313 as discussed above.

The position analyte sensor 296 is an analog reflecting light analyte sensor that has a light source and light receiver in the form of a photo transducer 317 disposed within a housing 318 with the housing 318 secured in fixed spatial relation to the driver coil pack 295. A reflective member 319 is disposed on or secured to a proximal end 320 of the magnetic member 301. The processor 313 determines the position of the lancet 299 by first emitting light from the light source of the photo transducer 317 towards the reflective member 319 with a predetermined solid angle of emission. Then, the light receiver of the photo transducer 317 measures the intensity of light reflected from the reflective member 319 and electrical conductors 315 transmit the signal generated there from to the processor 313.

By calibrating the intensity of reflected light from the reflective member 319 for various positions of the lancet 297 during the operating cycle of the driver coil pack 295, the position of the lancet 297 can thereafter be determined by measuring the intensity of reflected light at any given moment. In one embodiment, the analyte sensor 296 uses a commercially available LED/photo transducer module such as the OPB703 manufactured by Optek Technology, Inc., 1215 W. Crosby Road, Carrollton, Tex., 75006. This method of analog reflective measurement for position sensing can be used for any of the embodiments of lancet actuators discussed herein. In addition, any of the lancet actuators or drivers that include coils may use one or more of the coils to determine the position of the lancet 297 by using a magnetically permeable region on the lancet shaft 303 or magnetic member 301 itself as the core of a Linear Variable Differential Transformer

Referring to FIGS. 47 and 48, a flat coil lancet driver 325 is illustrated which has a main body housing 326 and a rotating frame 327. The rotating frame 327 pivots about an axle 328 disposed between a base 329, a top body portion 330 of the main body housing 326 and disposed in a pivot guide 331 of the rotating frame 327. An actuator arm 332 of the rotating frame 327 extends radially from the pivot guide 331 and has a linkage receiving opening 333 disposed at an outward end 334 of the actuator arm 332. A first end 335 of a coupler linkage 336 is coupled to the linkage receiving opening 333 of the actuator arm 332 and can rotate within the linkage receiving opening 333. A second end 337 of the coupler linkage 336 is disposed within an opening at a proximal end 338 of a coupler translation member 341. This configuration allows circumferential forces imposed upon the actuator arm 332 to be transferred into linear forces on a drive coupler 342 secured to a distal end 343 of the coupler translation member 341. The materials and dimensions of the drive coupler 342 can be the same or similar to the materials and dimensions of the drive coupler 342 discussed above.

Opposite the actuator arm 332 of the rotating frame 327, a translation substrate in the form of a coil arm 344 extends radially from the pivot guide 331 of the rotating frame 327. The coil arm 344 is substantially triangular in shape. A flat coil 345 is disposed on and secured to the coil arm 344. The flat coil 345 has leading segment 346 and a trailing segment 347, both of which extend substantially orthogonal to the direction of motion of the segments 346 and 347 when the rotating frame 327 is rotating about the pivot guide 331. The leading segment 346 is disposed within a first magnetically active region 348 generated by a first upper permanent magnet 349 secured to an upper magnet base 351 and a first lower permanent magnet 352 secured to a lower magnet base 353.

The trailing segment 347 is disposed within a second magnetically active region 354 generated by a second upper permanent magnet 355 secured to the upper magnet base 351 and a second lower permanent magnet secured to the lower magnet base 353.

The magnetic field lines or circuit of the first upper and lower permanent magnets 349, 352, 355 and 356 can be directed upward from the first lower permanent magnet 352 to the first upper permanent magnet 349 or downward in an opposite direction. The magnetic field lines from the second 10 permanent magnets 355 and 356 are also directed up or down, and will have a direction opposite to that of the first upper and lower permanent magnets 349 and 352. This configuration produces rotational force on the coil arm 344 about the pivot guide 331 with the direction of the force determined by the 15 direction of current flow in the flat coil 345.

A position analyte sensor 357 includes an optical encoder disk section 358 is secured to the rotating frame 327 which rotates with the rotating frame 327 and is read by an optical encoder 359 which is secured to the base 329. The position 20 analyte sensor 357 determines the rotational position of the rotating frame 327 and sends the position information to a processor 360 which can have features which are the same or similar to the features of the processor 193 discussed above via electrical leads 361. Electrical conductor leads 363 of the 25 flat coil 345 are also electrically coupled to the processor 360.

As electrical current is passed through the leading segment 346 and trailing segment 347 of the flat coil 345, the rotational forces imposed on the segments 346 and 347 are transferred to the rotating frame 327 to the actuator arm 332, through the 30 coupler linkage 336 and coupler translation member 341 and eventually to the drive coupler 342. In use, a lancet (not shown) is secured into the drive coupler 342, and the flat coil lancet actuator 325 activated. The electrical current in the flat coil 345 determines the forces generated on the drive coupler 35 342, and hence, a lancet secured to the coupler 342. The processor 360 controls the electrical current in the flat coil 345 based on the position and velocity of the lancet as measured by the position analyte sensor 357 information sent to the processor 360. The processor 360 is able to control the 40 velocity of a lancet in a manner similar to the processor 193 discussed above and can generate any of the desired lancet velocity profiles discussed above, in addition to others.

FIGS. 49 and 50 depict yet another embodiment of a controlled driver 369 having a driver coil pack 370 for a tissue 45 penetration device. The driver coil pack 370 has a proximal end 371, a distal end 372 and an axial lumen 373 extending from the proximal end 371 to the distal end 372. An inner coil 374 is disposed about the axial lumen 373 and has a tapered configuration with increasing wraps per inch of an elongate 50 conductor 375 in a distal direction. The inner coil 374 extends from the proximal end 371 of the coil driver pack 370 to the distal end 372 of the driver coil pack 370 with a major outer diameter or transverse dimension of about 1 to about 25 mm, specifically about 1 to about 12 mm.

The outer diameter or transverse dimension of the inner coil 374 at the proximal end 371 of the driver coil pack 370 is approximately equal to the diameter of the axial lumen 373 at the proximal end 371 of the coil pack 370. That is, the inner coil 374 tapers to a reduce outer diameter proximally until 60 there are few or no wraps of elongate electrical conductor 375 at the proximal end 371 of the driver coil pack 370. The tapered configuration of the inner coil 374 produces an axial magnetic field gradient within the axial lumen 373 of the driver coil pack 370 when the inner coil 374 is activated with 65 electrical current flowing through the elongate electrical conductor 375 of the inner coil 374.

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The axial magnetic field gradient produces a driving force for a magnetic member 376 disposed within the axial lumen 373 that drives the magnetic member 376 towards the distal end 372 of the driver coil pack 370 when the inner coil 374 is activated. The driving force on the magnetic member produced by the inner coil 374 is a smooth continuous force, which can produce a smooth and continuous acceleration of the magnetic member 376 and lancet 377 secured thereto. In some embodiments, the ratio of the increase in outer diameter versus axial displacement along the inner coil 374 in a distal direction can be from about 1 to about 0.08, specifically, about 1 to about 0.08.

An outer coil 378 is disposed on and longitudinally coextensive with the inner coil 374. The outer coil 378 can have the same or similar dimensions and construction as the inner coil 374, except that the outer coil 378 tapers proximally to an increased diameter or transverse dimension. The greater wraps per inch of elongate electrical conductor 379 in a proximal direction for the outer coil 378 produces a magnetic field gradient that drives the magnetic member 376 in a proximal direction when the outer coil 378 is activated with electrical current. This produces a braking or reversing effect on the magnetic member 376 during an operational cycle of the lancet 377 and driver coil pack 370. The elongate electrical conductors 375 and 379 of the inner coil 374 and outer coil 378 are coupled to a processor 381, which is coupled to an electrical power source 382. The processor 381 can have properties similar to the other processors discussed above and can control the velocity profile of the magnetic member 376 and lancet 377 to produce any of the velocity profiles above as well as others. The driver coil pack 370 can be used as a substitute for the coil driver pack discussed above, with other components of the lancing device 180 being the same or similar.

Embodiments of driver or actuator mechanisms having been described, we now discuss embodiments of devices which can house lancets, collect samples of fluids, analyze the samples or any combination of these functions. These front-end devices may be integrated with actuators, such as those discussed above, or any other suitable driver or controllable driver.

Generally, most known methods of blood sampling require several steps. First, a measurement session is set up by gathering various articles such as lancets, lancet drivers, test strips, analyzing instrument, etc. Second, the patient must assemble the paraphernalia by loading a sterile lancet, loading a test strip, and arming the lancet driver. Third, the patient must place a finger against the lancet driver and using the other hand to activate the driver. Fourth, the patient must put down the lancet driver and place the bleeding finger against a test strip, (which may or may not have been loaded into an analyzing instrument). The patient must insure blood has been loaded onto the test strip and the analyzing instrument has been calibrated prior to such loading. Finally, the patient must dispose of all the blood-contaminated paraphernalia including the lancet. As such, integrating the lancing and sample collection features of a tissue penetration sampling device can achieve advantages with regard to patient conve-

FIG. 51 shows a disposable sampling module 410, which houses the lancet 412. The lancet 412 has a head on a proximal end 416 which connects to the driver 438 and a distal end 414, which lances the skin. The distal end 414 is disposed within the conduit 418. The proximal end 416 extends into the cavity 420. The sample reservoir 422 has a narrow input port 424 on the ergonomically contoured surface 426, which is adjacent to the distal end 414 of the lancet 412. The term

ergonomically contoured, as used herein, generally means shaped to snugly fit a finger or other body portion to be lanced or otherwise tested placed on the surface. The sampling module **410** is capable of transporting the blood sample from the sample reservoir **422** through small passages (not shown), to 5 an analytical region **428**. The analytical region **428** can include chemical, physical, optical, electrical or other means of analyzing the blood sample. The lancet, sample flow channel, sample reservoir and analytical region are integrated into the sampling module **410** in a single packaged unit.

FIG. 52 shows the chamber 430 in the housing 410' where the sampling module 410 is loaded. The sampling module 410 is loaded on a socket 432 suspended with springs 434 and sits in slot 436. A driver 438 is attached to the socket 432. The driver 438 has a proximal end 440 and a distal end 442. The 15 driver 438 can be either a controllable driver or non-controllable driver any mechanical, such as spring or cam driven, or electrical, such as electromagnetically or electronically driven, means for advancing, stopping, and retracting the lancet. There is a clearance 444 between the distal end 442 of 20 the driver 438 and the analyte sensor 446, which is attached to the chamber 430. The socket 432 also contains an analyzer 448, which is a system for analyzing blood. The analyzer 448 corresponds to the analytical region 428 on the module 410 when it is loaded into the socket 432.

FIG. 53 shows a tissue penetration sampling device 411 with the sampling module 410 loaded into the socket 432 of housing 410'. The analytical region 428 and analyzer 448 overlap. The driver 438 fits into the cavity 420. The proximal end 440 of the driver 438 abuts the distal end 416 of the lancet 30 412. The patient's finger 450 sits on the ergonomically contoured surface 426.

FIG. **54** shows a drawing of an alternate lancet configuration where the lancet **412** and driver **438** are oriented to lance the side of the finger **450** as it sits on the ergonomically 35 contoured surface **426**.

FIG. 55 illustrates the orifice 452 and ergonomically contoured surface 426. The conduit 418 has an orifice 452, which opens on a blood well 454. The sample input port 424 of the reservoir 422 also opens on the blood well 454. The diameter 40 of the sample input port 424 is significantly greater than the diameter of the orifice 452, which is substantially the same diameter as the diameter of the lancet 412. After the lancet is retracted, the blood flowing from the finger 450 will collect in the blood well 454. The lancet 412 will have been retracted 45 into the orifice 452 effectively blocking the passage of blood down the orifice 452. The blood will flow from the blood well 454 through the sample input port 424 into the reservoir 422.

FIG. **56** shows a drawing of the lancing event. The patient applies pressure by pushing down with the finger 450 on the 50 ergonomically contoured surface 426. This applies downward pressure on the sampling module 410, which is loaded into the socket 432. As the socket 432 is pushed downward it compresses the springs 434. The analyte sensor 446 makes contact with the distal end 442 of the driver 438 and thereby 55 electrically detects the presence of the finger on the ergonomically contoured surface. The analyte sensor can be a piezoelectric device, which detects this pressure and sends a signal to circuit 456, which actuates the driver 438 and advances and then retracts the lancet 412 lancing the finger 60 450. In another embodiment, the analyte sensor 446 is an electric contact, which closes a circuit when it contacts the driver 438 activating the driver 438 to advance and retract the lancet 412 lancing the finger 450.

An embodiment of a method of sampling includes a 65 reduced number of steps that must be taken by a patient to obtain a sample and analysis of the sample. First, the patient

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loads a sampling module 410 with an embedded sterile lancet into the housing device 410'. Second, the patient initiates a lancing cycle by turning on the power to the device or by placing the finger to be lanced on the ergonomically contoured surface 426 and pressing down. Initiation of the analyte sensor makes the analyte sensor operational and gives control to activate the launcher.

The analyte sensor is unprompted when the lancet is retracted after its lancing cycle to avoid unintended multiple lancing events. The lancing cycle consists of arming, advancing, stopping and retracting the lancet, and collecting the blood sample in the reservoir. The cycle is complete once the blood sample has been collected in the reservoir. Third, the patient presses down on the sampling module, which forces the driver 38 to make contact with the analyte sensor, and activates the driver 438. The lancet then pierces the skin and the reservoir collects the blood sample.

The patient is then optionally informed to remove the finger by an audible signal such as a buzzer or a beeper, and/or a visual signal such as an LED or a display screen. The patient can then dispose of all the contaminated parts by removing the sampling module 410 and disposing of it. In another embodiment, multiple sampling modules 410 may be loaded into the housing 410' in the form of a cartridge (not shown). The patient can be informed by the tissue penetration sampling device 411 as to when to dispose of the entire cartridge after the analysis is complete.

In order to properly analyze a sample in the analytical region 428 of the sampling module 410, it may be desirable or necessary to determine whether a fluid sample is present in a given portion of the sample flow channel, sample reservoir or analytical area. A variety of devices and methods for determining the presence of a fluid in a region are discussed below.

In FIG. 57, a thermal analyte sensor 500 embedded in a substrate 502 adjacent to a surface 504 over which a fluid may flow. The surface may be, for example, a wall of a channel through which fluid may flow or a surface of a planar device over which fluid may flow. The thermal analyte sensor 500 is in electrical communication with a signal-conditioning element 506, which may be embedded in the substrate 502 or may be remotely located. The signal-conditioning element 506 receives the signal from the thermal analyte sensor 500 and modifies it by means such as amplifying it and filtering it to reduce noise. FIG. 57 also depicts a thermal analyte sensor 508 located at an alternate location on the surface where it is directly exposed to the fluid flow.

FIG. 58 shows a configuration of a thermal analyte sensor 500 adjacent to a separate heating element 510. The thermal analyte sensor 500 and the heating element 510 are embedded in a substrate 502 adjacent to a surface 504 over which a fluid may flow. In an alternate embodiment, one or more additional thermal analyte sensors may be adjacent the heating element and may provide for increased signal sensitivity. The thermal analyte sensor 500 is in electrical communication with a signal-conditioning element 506, which may be embedded in the substrate 502 or may be remotely located.

The signal-conditioning element 506 receives the signal from the thermal analyte sensor 500 and modifies it by means such as amplifying it and filtering it to reduce noise. The heating element 510 is in electrical communication with a power supply and control element 512, which may be embedded in the substrate 502 or may be remotely located. The power supply and control element 512 provides a controlled source of voltage and current to the heating element 510.

FIG. 59 depicts a configuration of thermal analyte sensors 500 having three thermal analyte sensor/heating element pairs (500/510), or detector elements, (with associated signal

conditioning elements 506 and power supply and control elements 512 as described in FIG. 58) embedded in a substrate 502 near each other alongside a surface 504. The figure depicts the thermal analyte sensors 500 arranged in a linear fashion parallel to the surface 504, but any operable configuration may be used. In alternate embodiments, fewer than three or more than three thermal analyte sensor/heating element pairs (500/510) may be used to indicate the arrival of fluid flowing across a surface 504. In other embodiments, self-heating thermal analyte sensors are used, eliminating the separate heating elements.

Embodiments of the present invention provide a simple and accurate methodology for detecting the arrival of a fluid at a defined location. Such detection can be particularly useful to define the zero- or start-time of a timing cycle for measuring rate-based reactions. This can be used in biochemical assays to detect a variety of analytes present in a variety of types of biological specimens or fluids and for rate-based reactions such as enzymatic reactions. Examples of relevant fluids include, blood, serum, plasma, urine, cerebral spinal fluid, saliva, enzymatic substances and other related substances and fluids that are well known in the analytical and biomedical art. The reaction chemistry for particular assays to analyze biomolecular fluids is generally well known, and 25 selection of the particular assay used will depend on the biological fluid of interest.

Assays that are relevant to embodiments of the present invention include those that result in the measurement of individual analytes or enzymes, e.g., glucose, lactate, creatinine kinase, etc, as well as those that measure a characteristic of the total sample, for example, clotting time (coagulation) or complement-dependent lysis. Other embodiments for this invention provide for sensing of sample addition to a test article or arrival of the sample at a particular location within that article.

Referring now to FIG. 60, a substrate 502 defines a channel 520 having an interior surface 522 over which fluid may flow. An analysis site 524 is located within the channel 520 where 40 fluid flowing in the channel 520 may contact the analysis site 524. In various embodiments, the analysis site 524 may alternatively be upon the interior surface 522, recessed into the substrate 502, or essentially flush with the interior surface 522. FIG. 60, depicts several possible locations for thermal 45 analyte sensors relative the substrate, the channel, and the analysis site; also, other locations may be useful and will depend upon the design of the device, as will be apparent to those of skill in art.

In use, thermal analyte sensors may be omitted from one or 50 more of the locations depicted in FIG. **60**, depending on the intended design. A recess in the analysis site **524** may provide the location for a thermal analyte sensor **526**, as may the perimeter of the analysis site provide the location for a thermal analyte sensor **528**. One or more thermal analyte sensors **530**, **532**, **534** may be located on the upstream side of the analysis site **524** (as fluid flows from right to left in FIG. **60**), or one or more thermal analyte sensors **536**, **538**, **540** may be located on the downstream side of the analysis site **524**.

The thermal analyte sensor may be embedded in the substrate near the surface, as thermal analyte sensor **542** is depicted. In various other embodiments, the thermal analyte sensor(s) may be located upon the interior surface, recessed into the interior surface, or essentially flush with the interior surface. Each thermal analyte sensor may also be associated 65 with a signal conditioning element, heating element, and power supply and control elements, as described above, and a

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single signal conditioning element, heating element, or power supply and control element may be associated with more than one thermal analyte sensor.

FIG. 61 shows possible positions for thermal analyte sensors relative to analysis sites 524 arranged in an array on a surface 556. A recess in the analysis site 524 may provide the location for a thermal analyte sensor 544, as may the perimeter of the analysis site provide the location for a thermal analyte sensor 546. The edge of the surface surrounding the array of analysis sites may provide the position for one or more thermal analyte sensors 548. Thermal analyte sensors may be positioned between analysis sites in a particular row 550 or column 552 of the array, or may be arranged on the diagonal 554.

In various embodiments, the thermal analyte sensor(s) may be may be embedded in the substrate near the surface or may be located upon the surface, recessed into the surface, or essentially flush with the surface. Each thermal analyte sensor may also be associated with a signal conditioning elements, heating elements, and power supply and control elements, as described above, and a single signal conditioning element, heating element, or power supply and control element may be associated with more than one thermal analyte sensor.

biomedical art. The reaction chemistry for particular assays to analyze biomolecular fluids is generally well known, and selection of the particular assay used will depend on the biological fluid of interest.

Assays that are relevant to embodiments of the present invention include those that result in the measurement of individual analytes or enzymes, e.g., glucose, lactate, creating invention and the proportion of the present individual analytes or enzymes, e.g., glucose, lactate, creating in the proportion of the present information about the biomolecular fluid being obtained through the use of reagents and/or test circuits and/or components associated with the analysis site.

FIG. 62 depicts several possible configurations of thermal analyte sensors relative to channels and analysis sites. The device schematically depicted in FIG. 62 may be, e.g., a microfluidic device for analyzing a small volume of a sample fluid, e.g. a biomolecular fluid. The device has a sample reservoir 560 for holding a quantity of a sample fluid. The sample fluid is introduced to the sample reservoir 560 via a sample inlet port 562 in fluid communication with the sample reservoir 560. A thermal analyte sensor 564 is located in or near the sample inlet port 562. A primary channel 566 originates at the sample reservoir 560 and terminates at an outflow reservoir 568.

One or more supplemental reservoirs 570 are optionally present and are in fluid communication with the primary channel 566 via one or more supplemental channels 572, which lead from the supplemental reservoir 570 to the primary channel 566. The supplemental reservoir 570 functions to hold fluids necessary for the operation of the assay, such as reagent solutions, wash solutions, developer solutions, fixative solutions, et cetera. In the primary channel 566 at a predetermined distance from the sample reservoir 560, an array of analysis sites 574 is present.

Thermal analyte sensors are located directly upstream (as fluid flows from right to left in the figure) from the array **576** and directly downstream from the array **578**. Thermal analyte sensors are also located in the primary channel adjacent to where the primary channel originates at the sample reservoir **580** and adjacent to where the primary channel terminates at the outflow reservoir **582**. The supplemental channel provides the location for another thermal analyte sensor **584**.

When the device is in operation, the thermal analyte sensor 564 located in or near the sample inlet port 562 is used to indicate the arrival of the sample fluid, e.g. the biomolecular fluid, in the local environment of the thermal analyte sensor, as described herein, and thus provides confirmation that the sample fluid has successfully been introduced into the device.

The thermal analyte sensor **580** located in the primary channel **566** adjacent to where the primary channel **566** originates at the sample reservoir **560** produces a signal indicating that sample fluid has started to flow from the sample reservoir **560** into the primary channel **566**. The thermal analyte sensors **576** in the primary channel **566** just upstream from the array of analysis sites **574** may be used to indicate that the fluid sample is approaching the array **574**. Similarly, the thermal analyte sensors **578** in the primary channel **566** just downstream from the array of analysis sites **574** may be used to indicate that the fluid sample has advanced beyond the array **574** and has thus contacted each analysis site.

The thermal analyte sensor **584** in the supplemental channel **572** provides confirmation that the fluid contained within the supplemental reservoir **570** has commenced to flow there from. The thermal analyte sensor **582** in the primary channel **566** adjacent to where the primary channel **566** terminates at the outflow reservoir **568** indicates when sample fluid arrives near the outflow reservoir **568**, which may then indicate that sufficient sample fluid has passed over the array of analysis sites **574** and that the analysis at the analysis sites is completed.

Embodiments of the invention provide for the use of a thermal analyte sensor to detect the arrival of the fluid sample 25 at a determined region, such as an analysis site, in the local environment of the thermal analyte sensor near the thermal analyte sensor. A variety of thermal analyte sensors may be used. Thermistors are thermally-sensitive resistors whose prime function is to detect a predictable and precise change in 30 electrical resistance when subjected to a corresponding change in temperature Negative Temperature Coefficient (NTC) thermistors exhibit a decrease in electrical resistance when subjected to an increase in temperature Coefficient (PTC) thermistors exhibit an 35 increase in electrical resistance when subjected to an increase in temperature.

A variety of thermistors have been manufactured for over the counter use and application. Thermistors are capable of operating over the temperature range of –100 degrees to over 40 600 degrees Fahrenheit. Because of their flexibility, thermistors are useful for application to micro-fluidics and temperature measurement and control.

A change in temperature results in a corresponding change in the electrical resistance of the thermistor. This temperature 45 change results from either an external transfer of heat via conduction or radiation from the sample or surrounding environment to the thermistor, or as an internal application of heat due to electrical power dissipation within the device. When a thermistor is operated in "self-heating" mode, the power dissipated in the device is sufficient to raise its temperature above the temperature of the local environment, which in turn more easily detects thermal changes in the conductivity of the local environment.

Thermistors are frequently used in "self heating" mode in 55 applications such as fluid level detection, airflow detection and thermal conductivity materials characterization. This mode is particularly useful in fluid sensing, since a self-heating conductivity analyte sensor dissipates significantly more heat in a fluid or in a moving air stream than it does in 60 still air.

Embodiments of the invention may be designed such that the thermal analyte sensor is exposed directly to the sample. However, it may also be embedded in the material of the device, e.g., in the wall of a channel meant to transport the 65 sample. The thermal analyte sensor may be covered with a thin coating of polymer or other protective material.

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Embodiments of the device need to establish a baseline or threshold value of a monitored parameter such as temperature. Ideally this is established during the setup process. Once fluid movement has been initiated, the device continuously monitors for a significant change thereafter. The change level designated as "significant" is designed as a compromise between noise rejection and adequate sensitivity. The actual definition of the "zero- or start-time" may also include an algorithm determined from the time history of the data, i.e., it can be defined ranging from the exact instant that a simple threshold is crossed, to a complex mathematical function based upon a time sequence of data.

In use, a signal is read from a thermal analyte sensor in the absence of the sample or fluid. The fluid sample is then introduced. The sample flows to or past the site of interest in the local environment of the thermal analyte sensor, and the thermal analyte sensor registers the arrival of the sample. The site of interest may include an analysis site for conducting, e.g., an enzymatic assay. Measuring the arrival of fluid at the site of interest thus indicates the zero- or start-time of the reaction to be performed. For detection of fluid presence, these sites may be any of a variety of desired locations along the fluidic pathway. Embodiments of the invention are particularly well suited to a microfluidic cartridge or platform, which provide the user with an assurance that a fluid sample has been introduced and has flowed to the appropriate locations in the platform.

A rate-based assay must measure both an initiation time, and some number of later time points, one of which is the end-point of the assay. Therefore, baseline or threshold value can be established, and then continuously monitored for a significant change thereafter; one such change is the arrival of the fluid sample that initiates the enzyme reaction. Baseline values are frequently established during the device setup process. The threshold is designed as a compromise between noise rejection and adequate sensitivity. The defined zero- or "start-time" can be defined ranging from the exact instant that a simple threshold is crossed, to the value algorithmically determined using a filter based on a time sequence of data.

Embodiments of the invention accomplish this in a variety of ways. In one embodiment, an initial temperature measurement is made at a thermal analyte sensor without the sample present. The arrival of a sample changes causes the thermal analyte sensor to register a new value. These values are then compared.

Another embodiment measures the change in thermal properties (such as thermal conductivity or thermal capacity) in the local environment of a thermal analyte sensor caused by the arrival of a fluid sample. In general this is the operating principle of a class of devices known as "thermal conductivity analyte sensors" or "heat flux analyte sensors". At least two hardware implementations have been used and are described above. One implementation utilizes a thermal analyte sensor in a "self-heating mode." In "self-heating mode," a self-heating thermal analyte sensor may utilize a positive temperature coefficient thermistor placed in or near the flow channel, e.g. located in the wall of the flow channel.

An electrical current is run through the thermistor, causing the average temperature of the thermistor to rise above that of the surrounding environment. The temperature can be determined from the electrical resistance, since it is temperature dependent. When fluid flows through the channel, it changes the local thermal conductivity near the thermistor (usually to become higher) and this causes a change in the average temperature of the thermistor. It also changes the thermal capacity, which modifies the thermal dynamic response. These

changes give rise to a signal, which can be detected electronically by well-known means, and the arrival of the fluid can thereby be inferred.

A second hardware implementation requires a separate heating element in or near the flow channel, plus a thermal 5 analyte sensor arrangement in close proximity. Passing a current through the element provides heat to the local environment and establishes a local temperature detected by the thermocouple device. This temperature or its dynamic response is altered by the arrival of the fluid or blood in or near 10 the local environment, similar to the previously described implementation, and the event is detected electronically.

The heating element can be operated in a controlled input mode, which may include controlling one or more of the following parameters-applied current, voltage or power—in a 15 prescribed manner. When operating in controlled input mode, fluctuations of the temperature of the thermal analyte sensor are monitored in order to detect the arrival of the fluid.

Alternatively, the heating element can be operated in such a fashion as to control the temperature of the thermal analyte 20 sensor in a prescribed manner. In this mode of operation, the resulting fluctuations in one or more of the input parameters to the heating element (applied current, voltage, and power) can be monitored in order to detect the arrival of the fluid.

In either of the above-described operating modes, the prescribed parameter can be held to a constant value or sequence of values that are held constant during specific phases of operation of the device. The prescribed parameter can also varied as a known function or waveform in time.

The change in the monitored parameters caused by the 30 arrival of the fluid can be calculated in any of a number of ways, using methods well known in the art of signal processing. The signal processing methods allow the relation of the signal received prior to arrival of the fluid with the signal received upon arrival of the fluid to indicate that the fluid has 35 arrived. For example, and after suitable signal filtering is applied, changes in the monitored value or the rate of change of the value of the signal can be monitored to detect the arrival of the fluid. Additionally, the arrival of fluid will cause a dynamic change in the thermodynamic properties of the local 40 environment, such as thermal conductivity or thermal capacity. When the input parameter is a time varying function this change of thermodynamic properties will cause a phase shift of the measured parameter relative to the controlled parameter. This phase shift can be monitored to detect the arrival of 45 the fluid.

It should also be noted that sensitivity to thermal noise and operating power levels could be reduced in these either of these modes of operation by a suitable choice of time-varying waveforms for the prescribed parameter, together with appropriate and well-known signal processing methods applied to the monitored parameters. However, these potential benefits may come at the cost of slower response time.

Referring to FIG. 63, an alternative embodiment of a tissue penetration sampling device is shown which incorporates 55 disposable sampling module 590, a lancet driver 591, and an optional module cartridge 592 are shown. The optional module cartridge comprises a case body 593 having a storage cavity 594 for storing sampling modules 590. A cover to this cavity has been left out for clarity. The cartridge further comprises a chamber 595 for holding the lancet driver 591. The lancet driver has a preload adjustment knob 596, by which the trigger point of the lancet driver may be adjusted. This insures a reproducible tension on the surface of the skin for better control of the depth of penetration and blood yield. 65 In one embodiment, the sampling module 590 is removably attached to the lancet driver 591, as shown, so that the sam-

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pling module **590** is disposable and the lancet driver **591** is reusable. In an alternative embodiment, the sampling module and lancet driver are contained within a single combined housing, and the combination sample acquisition module/lancet driver is disposable. The sampling module **590** includes a sampling site **597**, preferably having a concave depression **598**, or cradle, that can be ergonomically designed to conform to the shape of a user's finger or other anatomical feature (not shown).

The sampling site further includes an opening 599 located in the concave depression. The lancet driver 591 is used to fire a lancet contained within and guided by the sampling module 590 to create an incision on the user's finger when the finger is placed on the sampling site 597. In one embodiment, the sampling site forms a substantially airtight seal at the opening when the skin is firmly pressed against the sampling site; the sampling site may additionally have a soft, compressible material surrounding the opening to further limit contamination of the blood sample by ambient air. "Substantially airtight" in this context means that only a negligible amount of ambient air may leak past the seal under ordinary operating conditions, the substantially airtight seal allowing the blood to be collected seamlessly.

Referring to FIGS. **64** and **65**, the lancet **600** is protected in the integrated housing **601** that provides a cradle **602** for positioning the user's finger or other body part, a sampling port **603** within the cradle **602**, and a sample reservoir **603**' for collecting the resulting blood sample. The lancet **600** is a shaft with a distal end **604** sharpened to produce the incision with minimal pain. The lancet **600** further has an enlarged proximal end **605** opposite the distal end. Similar lancets are commonly known in the art.

Rather than being limited to a shaft having a sharp end, the lancet may have a variety of configurations known in the art, with suitable modifications being made to the system to accommodate such other lancet configurations, such configurations having a sharp instrument that exits the sampling port to create a wound from which a blood sample may be obtained.

In the figures, the lancet 600 is slidably disposed within a lancet guide 606 in the housing 601, and movement of the lancet 600 within the lancet guide 606 is closely controlled to reduce lateral motion of the lancet, thereby reducing the pain of the lance stick. The sample acquisition module also includes a return stop 613, which retains the lancet within the sample acquisition module. The sampling module has an attachment site 615 for attachment to the lancet driver.

The sampling module further includes a depth selector allowing the user to select one of several penetration depth settings. The depth selector is shown as a multi-position thumbwheel 607 having a graduated surface. By rotating the thumbwheel 607, the user selects which part of the graduated surface contacts the enlarged proximal end 605 of the lancet to limit the movement of the lancet 600 within the lancet guide 606.

The thumbwheel is maintained in the selected position by a retainer 608 having a protruding, rounded surface which engages at least one of several depressions 609 (e.g. dimples, grooves, or slots) in the thumbwheel 607. The depressions 609 are spatially aligned to correspond with the graduated slope of the thumbwheel 607, so that, when the thumbwheel 607 is turned, the depth setting is selected and maintained by the retainer 608 engaging the depression 609 corresponding to the particular depth setting selected.

In alternate embodiments, the retainer may be located on the depth selector and the depressions corresponding to the depth setting located on the housing such that retainer may

functionally engage the depressions. Other similar arrangements for maintaining components in alignment are known in the art and may be used. In further alternate embodiments, the depth selector may take the form of a wedge having a graduated slope, which contacts the enlarged proximal end of the lancet, with the wedge being retained by a groove in the housing.

The sample reservoir **603'** includes an elongate, rounded chamber **610** within the housing **601** of the sample acquisition module. The chamber **610** has a flat or slightly spherical shape, with at least one side of the chamber **610** being formed by a smooth polymer, preferably absent of sharp corners. The sample reservoir **603'** also includes a sample input port **611** to the chamber **610**, which is in fluid communication with the sampling port **603**, and a vent **612** exiting the chamber.

A cover (not shown), preferably of clear material such as plastic, positions the lancet 600 and closes the chamber 603', forming an opposing side of the chamber 603'. In embodiments where the cover is clear, the cover may serve as a testing means whereby the sample may be analyzed in the 20 reservoir via optical sensing techniques operating through the cover. A clear cover will also aid in determining by inspection when the sample reservoir is full of the blood sample.

FIG. 66 shows a portion of the sampling module illustrating an alternate embodiment of the sample reservoir. The 25 sample reservoir has a chamber 616 having a sample input port 617 joining the chamber 616 to a blood transport capillary channel 618; the chamber 616 also has a vent 619. The chamber has a first side 620 that has a flat or slightly spherical shape absent of sharp corners and is formed by a smooth 30 polymer. An elastomeric diaphragm 621 is attached to the perimeter of the chamber 616 and preferably is capable of closely fitting to the first side of the chamber 620.

To control direction of blood flow, the sample reservoir is provided with a first check valve 622 located at the entrance 35 617 of the sample reservoir and a second check valve 623 leading to an exit channel 624 located at the vent 619. Alternately, a single check valve (at the location 622) may be present controlling both flow into the chamber 616 via the blood transport capillary channel 618 and flow out of the 40 chamber 616 into an optional alternate exit channel 625. The sample reservoir has a duct 626 connecting to a source of variable pressure facilitating movement of the diaphragm

When the diaphragm 621 is flexed away from the first side 45 of the chamber 620 (low pressure supplied from the source via duct 626), the first check valve 622 is open and the second check valve 623 is closed, aspiration of the blood sample into the sample reservoir follows. When the diaphragm 621 is flexed in the direction of the first side of the chamber 620 (high pressure supplied from the source via duct 626) with the first check valve 622 closed and the second check valve 623 open, the blood is forced out of the chamber 616. The direction of movement and actuation speed of the diaphragm 621 can be controlled by the pressure source, and therefore the 55 flow of the sample can be accelerated or decelerated. This feature allows not only reduced damage to the blood cells but also for the control of the speed by which the chamber 616 is filled.

While control of the diaphragm 621 via pneumatic means 60 is described in this embodiment, mechanical means may alternately be used. Essentially, this micro diaphragm pump fulfills the aspiration, storage, and delivery functions. The diaphragm 621 may be used essentially as a pump to facilitate transfer of the blood to reach all areas required. Such required 65 areas might be simple sample storage areas further downstream for assaying or for exposing the blood to a chemical

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analyte sensor or other testing means. Delivery of the blood may be to sites within the sampling module or to sites outside the sampling module, i.e. a separate analysis device.

In an alternate embodiment, a chemical analyte sensor or other testing means is located within the sampling module, and the blood is delivered to the chemical analyte sensor or other testing means via a blood transfer channel in fluid communication with the sample reservoir. The components of the sampling module may be injection molded and the diaphragm may be fused or insertion molded as an integral component.

FIG. 67 depicts a portion of the disposable sampling module surrounding the sampling port 627, including a portion of the sampling site cradle surface 628. The housing of the sampling module includes a primary sample flow channel 629 that is a capillary channel connecting the sample input port to the sample reservoir. The primary sample flow channel 629 includes a primary channel lumenal surface 630 and a primary channel entrance 631, the primary channel entrance 631 opening into the sample input port 627. The sampling module may optionally include a supplemental sample flow channel 632 that is also a capillary channel having a supplemental channel lumenal surface 633 and a supplemental channel entrance 634, the supplemental channel entrance 634 opening into the sample input port 627.

The primary sample flow channel 629 has a greater crosssectional area than the supplemental sample flow channel 632, preferably by at least a factor of two. Thus, the supplemental sample flow channel 632 draws fluid faster than the primary sample flow channel 629. When the first droplet of blood is received into the sample input port 627, the majority of this droplet is drawn through the supplemental sample flow channel 632. However, as the blood continues to flow from the incision into the sample input port 627, most of this blood is drawn through the primary sample flow channel 629, since the supplemental sample flow channel 632 is of limited capacity and is filled or mostly filled with the first blood droplet. This dual capillary channel configuration is particularly useful in testing where there is a concern with contamination of the sample, e.g. with debris from the lancet strike or (particularly in the case of blood gas testing) with air.

In order to improve blood droplet flow, some priming or wicking of the surface with blood is at times necessary to begin the capillary flow process. Portions of the surfaces of the sample input port 627 and the primary and supplemental (if present) sample flow channels 629, 632 are treated to render those surfaces hydrophilic. The surface modification may be achieved using mechanical, chemical, corona, or plasma treatment. Examples of such coatings and methods are marketed by AST Products (Billerica, Mass.) and Spire Corporation (Bedford, Mass.).

However, a complete blanket treatment of the surface could prove detrimental by causing blood to indiscriminately flow all over the surface and not preferentially through the capillary channel(s). This ultimately will result in losses of blood fluid. The particular surfaces which receive the treatment are selected to improve flow of blood from an incised finger on the sampling site cradle surface 628 through the sample input port 627 and at least one of the sample flow channels 629, 632 to the sample reservoir. Thus, the treatment process should be masked off and limited only to the selected surfaces. The masking process of selectively modifying the sampling surface from hydrophobic to hydrophilic may be done with mechanical masking techniques such as with metal shielding, deposited dielectric or conductive films, or electrical shielding means.

In some embodiments, the treated surfaces are limited to one or more of the following: the surface of the sampling port which lies between the sampling site cradle surface and the primary and supplemental sample flow channel, the surface immediately adjacent to the entrances to the primary and/or supplemental sample flow channels 631, 634 (both within the sample input port and within the sample flow channel), and the lumenal surface of the primary and/or supplemental sample flow channels 630, 633.

Upon exiting the incision blood preferentially moves 10 through the sample input port 627 into the supplementary sample flow channel 632 (if present) and into the primary sample flow channel 629 to the sample reservoir, resulting in efficient capture of the blood. Alternatively, the substrate material may be selected to be hydrophilic or hydrophobic, 15 and a portion of the surface of the substrate material may be treated for the opposite characteristic.

In an embodiment, FIG. 67 a membrane 635 at the base of the sample input port 627 is positioned between the retracted sharpened distal end of the lancet 636 and the entrance to the sample flow channels 631, 634. The membrane 635 facilitates the blood sample flow through the sample flow channels 629, 632 by restricting the blood from flowing into the area 636 surrounding the distal end of the lancet 637. The blood thus flows preferentially into the sample reservoir. In an embodiment, the membrane 635 is treated to have a hydrophobic characteristic. In another embodiment, the membrane 635 is made of polymer-based film 638 that has been coated with a silicone-based gel 639.

For example, the membrane structure may comprise a 30 polymer-based film 638 composed of polyethylene terephthalate, such as the film sold under the trademark MYLAR. The membrane structure may further comprise a thin coating of a silicone-based gel 639 such as the gel sold under the trademark SYLGARD on at least one surface of the film. The usefulness of such a film is its ability to reseal after the lancet has penetrated it without physically affecting the lancet's cutting tip and edges. The MYLAR film provides structural stability while the thin SYLGARD silicone laminate is flexible enough to retain its form and close over the hole made in the MYLAR film. Other similar materials fulfilling the structural stability and flexibility roles may be used in the manufacture of the membrane in this embodiment.

The membrane 635 operates to allow the sharpened distal end of the lancet 637 to pierce the membrane as the sharpened 45 distal end of the lancet 637 travels into and through the sample input port 627. In an embodiment, the silicone-based gel 639 of the membrane 635 automatically seals the cut caused by the piercing lancet. Therefore, after an incision is made on a finger of a user, the blood from the incision is prevented from 50 flowing through the membrane 635, which aids the blood to travel through the primary sample flow channel 629 to accumulate within the sample reservoir. Thus the film prevents any blood from flowing into the lancet device assembly, and blood contamination and loss into the lancet device mechanism 55 cavity are prevented. Even without the resealing layer 639, the hydrophobic membrane 635 deters the flow of blood across the membrane 635, resulting in improved flow through the primary sample flow channel 629 and reduced or eliminated flow through the pierced membrane 635.

FIGS. **68-70** illustrate one implementation of a lancet driver **640** at three different points during the use of the lancet driver. In this description of the lancet driver, proximal indicates a position relatively close to the site of attachment of the sampling module; conversely, distal indicates a position relatively far from the site of attachment of the sampling module. The lancet driver has a driver handle body **641** defining a

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cylindrical well 642 within which is a preload spring 643. Proximal to the preload spring 643 is a driver sleeve 644, which closely fits within and is slidably disposed within the well 642. The driver sleeve 644 defines a cylindrical driver chamber 645 within which is an actuator spring 646. Proximal to the actuator spring 646 is a plunger sleeve 647, which closely fits within and is slidably disposed within the driver sleeve 644.

The driver handle body **641** has a distal end **648** defining a threaded passage **649** into which a preload screw **650** fits. The preload screw defines a counterbore **651**. The preload screw **650** has a distal end **652** attached to a preload adjustment knob **653** and a proximal end **654** defining an aperture **655**. The driver sleeve **644** has a distal end **656** attached to a catch fitting **657**. The catch fitting **657** defines a catch hole **658**. The driver sleeve **644** has a proximal end **659** with a sloped ring feature **660** circling the interior surface of the driver sleeve's proximal end **659**.

The lancet driver includes a plunger stem 660 having a proximal end 661 and a distal end 662. At its distal end 662. an enlarged plunger head 663 terminates the plunger stem 660. At its proximal end 661, the plunger stem 660 is fixed to the plunger tip 667 by adhesively bonding, welding, crimping, or threading into a hole 665 in the plunger tip 667. A plunger hook 665 is located on the plunger stem 660 between the plunger head 663 and the plunger tip 667. The plunger head 663 is slidably disposed within the counterbore 651 defined by the preload screw 650. The plunger stem 660 extends from the plunger head 663, through the aperture 655 defined by the proximal end 654 of the preload screw, thence through the hole 658 in the catch fitting 657, to the joint 664 in the plunger tip 667. For assembly purposes, the plunger base joint 664 may be incorporated into the plunger sleeve 647, and the plunger stem 660 attached to the plunger base 664 by crimping, swaging, gluing, welding, or some other means. Note that the lancet driver 640 could be replaced with any of the controlled electromagnetic drivers discussed above.

stability while the thin SYLGARD silicone laminate is flexible enough to retain its form and close over the hole made in the MYLAR film. Other similar materials fulfilling the structural stability and flexibility roles may be used in the manufacture of the membrane in this embodiment.

The membrane 635 operates to allow the sharpened distalend of the lancet 637 to pierce the membrane as the sharpened distalend of the lancet 637 travels into and through the sample input port 627. In an embodiment, the silicone-based gel 639 of the membrane 635 automatically seals the cut caused by

The combined sampling module and lancet driver assembly is then pressed against the user's finger (or other selected anatomical feature) in a smooth motion until the preset trigger point is reached. The trigger point corresponds to the amount of preload force that needs to be overcome to actuate the driver to drive the lancet towards the skin. The preload screw allows the preload setting to be adjusted by the user such that a consistent, preset (by the user) amount of preload force is applied to the sampling site 597 each time a lancing is performed.

When the motion to press the assembly against the user's finger is begun (see FIG. 68), the plunger hook 665 engages catch fitting 657, holding the actuator spring 646 in a cocked position while the force against the finger builds as the driver sleeve 644 continues to compress the preload spring 643. Eventually (see FIG. 69) the sloped back of the plunger hook 665 slides into the hole 655 in the proximal end of the preload screw 654 and disengages from the catch fitting 657. The plunger sleeve 647 is free to move in a proximal direction

once the plunger hook 665 releases, and the plunger sleeve 647 is accelerated by the actuator spring 646 until the plunger tip 667 strikes the enlarged proximal end of the lancet 212.

Upon striking the enlarged proximal end of the lancet **605**, the plunger tip **667** of the actuated lancet driver reversibly 5 engages the enlarged proximal end of the lancet **605**. This may be accomplished by mechanical means, e.g. a fitting attached to the plunger tip **667** that detachably engages a complementary fitting on the enlarged proximal end of the lancet **605**, or the enlarged proximal end of the lancet **605** may 10 be coated with an adhesive that adheres to the plunger tip **667** of the actuated lancet driver. Upon being engaged by the plunger tip **667**, the lancet **600** slides within the lancet guide **606** with the sharpened distal end of the lancet **604** emerging from the housing **601** through the sampling port **603** to create 15 the incision in the user's finger.

At approximately the point where the plunger tip 667 contacts the enlarged proximal end of the lancet 605, the actuator spring 646 is at its relaxed position, and the plunger tip 667 is traveling at its maximum velocity. During the extension 20 stroke, the actuator spring 646 is being extended and is slowing the plunger tip 667 and lancet 600. The end of stroke occurs (see FIG. 70) when the enlarged proximal end of the lancet 605 strikes the multi-position thumbwheel 607.

The direction of movement of the lancet **600** is then 25 reversed and the extended actuator spring then quickly retracts the sharpened distal end of the lancet **604** back through the sampling port **603**. At the end of the return stroke, the lancet **600** is stripped from the plunger tip **667** by the return stop **613**. The adhesive adheres to the return stop **613** 30 retaining the lancet in a safe position.

As blood seeps from the wound, it fills the sample input port 603 and is drawn by capillary action into the sample reservoir 603'. In this embodiment, there is no reduced pressure or vacuum at the wound, i.e. the wound is at ambient air 35 pressure, although embodiments which draw the blood sample by suction, e.g. supplied by a syringe or pump, may be used. The vent 612 allows the capillary action to proceed until the entire chamber is filled, and provides a transfer port for analysis of the blood by other instrumentation. The finger is 40 held against the sample acquisition module until a complete sample is observed in the sample reservoir.

As the sampling module 600 is removed from the lancet driver 591, a latch 614 that is part of the return stop 613 structure engages a sloped ring feature 660 inside the lancet 45 driver 591. As the lancet driver 591 is removed from the sampling module 600, the latch forces the return stop 613 to rotate toward the lancet 600, bending it to lock it in a safe position, and preventing reuse.

As the sampling module 600 is removed from the lancet 50 driver 591, the driver sleeve 644 is forced to slide in the driver handle body 641 by energy stored in the preload spring 643. The driver sleeve 644, plunger sleeve 647, and actuator spring 646 move outward together until the plunger head 663 on the plunger stem 660 contacts the bottom of the counterbore 651 at the proximal end of the preload screw 654. The preload spring 643 continues to move the driver sleeve 644 outward compressing the actuator spring 646 until the plunger hook 665 passes through the hole 658 in the catch fitting 657. Eventually the two springs reach equilibrium and the plunger 60 sleeve 647 comes to rest in a cocked position.

After the sampling module 600 is removed from the lancet driver 591 it may be placed in a separate analysis device to obtain blood chemistry readings. In a preferred embodiment, the integrated housing 601 or sample reservoir 603' of the 65 sampling module 600 contains at least one bioanalyte sensor, which is powered by and/or read by the separate analysis

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device. In another embodiment, the analysis device performs an optical analysis of the blood sample directly through the clear plastic cover of the sampling module. Alternatively, the blood sample may be transferred from the sampling module into an analysis device for distribution to various analysis processes.

Alternate embodiments of the invention offer improved success rates for sampling, which reduces the needless sacrifice of a sample storage reservoir or an analysis module due to inadequate volume fill. Alternate embodiments allow automatic verification that sufficient blood has been collected before signaling the user (e.g. by a signal light or an audible beep) that it is okay to remove the skin from the sampling site. In such alternate embodiments, one or more additional lancet (s) (denoted backup lancets) and/or lancet driver(s) (denoted backup lancet drivers) and/or sample reservoir(s) (denoted backup sample reservoirs) are present with the "primary" sampling module.

In one such preferred embodiment, following detection of inadequate blood sample volume (e.g., by light or electronic methods), a backup sampling cycle is initiated automatically. The "backup sampling cycle" includes disconnecting the primary sample reservoir via a simple valving system, bringing the backup components online, lancing of the skin, collection of the blood, and movement of the blood to the backup sample reservoir.

Blood flows into the backup sample reservoir until the required volume is obtained. The cycle repeats itself, if necessary, until the correct volume is obtained. Only then is the sample reservoir made available as a source of sampled blood for use in measurements or for other applications. The series of reservoirs and/or lancets and/or lancet drivers may easily be manufactured in the same housing and be transparent to the user.

In one embodiment, up to three sample reservoirs (the primary plus two backup) are present in a single sample acquisition module, each connected via a capillary channel/valving system to one or more sampling ports. Another embodiment has four sample reservoirs (the primary plus three backup) present in a single sample acquisition module, each connected via a capillary channel/valving system to one or more sampling ports. With three or four sample reservoirs, at least an 80% sampling success rate can be achieved for some embodiments.

Another embodiment includes a miniaturized version of the tissue penetration sampling device. Several of the miniature lancets may be located in a single sampling site, with corresponding sample flow channels to transfer blood to one or more reservoirs. The sample flow channels may optionally have valves for controlling flow of blood. The device may also include one or more analyte sensors, such as the thermal analyte sensors discussed above, for detecting the presence of blood, e.g. to determine if a sufficient quantity of blood has been obtained. In such an embodiment, the disposable sampling module, the lancet driver, and the optional module cartridge will have dimensions no larger than about 150 mm long, 60 mm wide, and 25 mm thick.

In other embodiments, the size of the tissue penetration sampling device including the disposable sampling module, the lancet driver, and the optional cartridge will have dimensions no larger than about 100 mm long, about 50 mm wide, and about 20 mm thick, and in still other embodiments no larger than about 70 mm long, about 30 mm wide, and about 10 mm thick. The size of the tissue penetration sampling device including the disposable sampling module, the lancet driver, and the optional cartridge will generally be at least about 10 mm long, about 5 mm wide, and about 2 mm thick.

In another miniature embodiment, the dimensions of the lancet driver without the cartridge or sampling module are no larger than about 80 mm long, 10 mm wide, and 10 mm thick, or specifically no larger than about 50 mm long, 7 mm wide, and 7 mm thick, or even more specifically no larger than about 5 mm long, 5 mm wide, and 3 mm thick; dimensions of the lancet driver without the cartridge or sampling module are generally at least about 1 mm long, 0.1 mm wide, and 0.1 mm thick, or specifically at least about 2 mm long, 0.2 mm wide, and 0.2 mm thick, or more specifically at least about 4 mm 10 long, 0.4 mm wide, and 0.4 mm thick.

In yet another miniature embodiment, dimensions of the miniature sampling module without the lancet driver or cartridge are no larger than about 15 mm long, about 10 mm wide, and about 10 mm thick, or no larger than about 10 mm thick, or no larger than about 5 mm long, about 3 mm wide, and about 2 mm thick; dimensions of the miniature sampling module without the lancet driver or cartridge are generally at least about 1 mm long, 0.1 mm wide, and 0.1 mm thick, specifically at least about 2 mm long, 0.2 mm wide, and 0.2 mm thick, or more specifically at least about 4 mm long, 0.4 mm wide, and 0.4 mm thick.

In another embodiment, the miniaturized sampling module and the lancet driver form a single unit having a shared housing, and the combined sample acquisition module/lancet driver unit is disposable. Such a combined unit is no larger than about 80 mm long, about 30 mm wide, and about 10 mm thick, specifically no larger than about 50 mm long, about 20 mm wide, and about 5 mm thick, more specifically, no larger than about 20 mm long, about 5 mm wide, and about 3 mm thick; the combined unit is generally at least about 2 mm long, about 0.3 mm wide, and about 0.2 mm thick, specifically at least about 4 mm long, 0.6 mm wide, and 0.4 mm thick, more specifically, at least about 8 mm long, 1 mm wide, and 0.8 mm 35 thick

Referring to FIG. 71, another embodiment of a tissue penetration sampling device is shown, incorporating a disposable sampling module 608 cartridge and analyzer device 669 is shown. The analyzer device 669 includes a deck 670 having a 40 lid 671 attached to the deck by hinges along the rear edge of the system 672. A readout display 673 on the lid 671 functions to give the user information about the status of the analyzer device 669 and/or the sampling module cartridge 668, or to give readout of a blood test. The analyzer device 669 has several function buttons 674 for controlling function of the analyzer device 669. Alternatively, the reader device may have a touch-sensitive screen, an optical scanner, or other input means known in the art.

An analyzer device with an optical scanner may be particularly useful in a clinical setting, where patient information may be recorded using scan codes on patients' wristbands or files. The analyzer reader device may have a memory, enabling the analyzer device to store results of many recent tests. The analyzer device may also have a clock and calendar function, enabling the results of tests stored in the memory to be time and date-stamped. A computer interface 675 enables records in memory to be exported to a computer. The analyzer device 669 has a chamber located between the deck 670 and 60 the lid 671, which closely accommodates a sampling module cartridge 668. Raising the lid 671, allowing a sampling module cartridge 668 to be inserted or removed, accesses the chamber.

FIG. 72 is an illustration showing some of the features of an 65 embodiment of a sampling module cartridge. The sampling module cartridge 668 has a housing having an orientation

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sensitive contact interface for mating with a complementary surface on the analyzer device. The contact interface functions to align the sampling module cartridge with the analyzer device, and also allows the analyzer device to rotate the sampling module cartridge in preparation for a new sampling event. The contact interface may take the form of cogs or grooves formed in the housing, which mate with complementary cogs, or grooves in the chamber of the analyzer device.

The sampling module cartridge has a plurality of sampling sites 678 on the housing, which are shown as slightly concave depressions near the perimeter of the sampling module cartridge 668. Each sampling site defines an opening 679 contiguous with a sample input port entering the sampling module. In an alternate embodiment, the sampling sites and sample input ports are located on the edge of the sampling module cartridge. Optical windows 680 allow transmission of light into the sampling module cartridge for the purpose of optically reading test results. Alternatively, analyte sensor connection points allow transmission of test results to the analyzer device via electrical contact. Access ports 681, if present, allow transmission of force or pressure into the sampling module cartridge from the analyzer device. The access ports may be useful in conjunction with running a calibration test or combining reagents with sampled blood or other bodily fluids.

The described features are arranged around the sampling module cartridge, and the sampling module cartridge is radially partitioned into many sampling modules, each sampling module having the components necessary to perform a single blood sampling and testing event. A plurality of sampling modules are present on a sampling module cartridge, generally at least ten sampling modules are present on a single disposable sampling module cartridge; at least about 20, or more on some embodiments, and at least about 34 sampling modules are present on one embodiment, allowing the sampling module cartridge to be maintained in the analyzer device for about a week before replacing with a new sampling module cartridge (assuming five sampling and testing events per day for seven days). With increasing miniaturization, up to about 100, or more preferably up to about 150, sampling modules may be included on a single sampling module cartridge, allowing up to a month between replacements with new sampling module cartridges. It may be necessary for sampling sites to be located in several concentric rings around the sampling module cartridge or otherwise packed onto the housing surface to allow the higher number of sampling modules on a single sampling module cartridge.

In other embodiments, the sampling module cartridge may be any other shape which may conveniently be inserted into a analyzer device and which are designed to contain multiple sampling modules, e.g. a square, rectangular, oval, or polygonal shape. Each sampling module is miniaturized, being generally less than about 6.0 cm long by about 1.0 cm wide by about 1.0 cm thick, so that thirty five more or less wedgeshaped sampling modules can fit around a disk having a radius of about 6.0 cm. In some embodiments, the sampling modules can be much smaller, e.g. less than about 3.0 cm long by about 0.5 cm wide by about 0.5 cm thick.

FIG. 73 depicts, in a highly schematic way, a single sampling module, positioned within the analyzer device. Of course, it will occur to the person of ordinary skill in the art that the various recited components may be physically arranged in various configurations to yield a functional system. FIG. 73 depicts some components, which might only be present in alternate embodiments and are not necessarily all present in any single embodiment. The sampling module has a sample input port 682, which is contiguous with an opening

**683** defined by a sampling site **684** on the cartridge housing **685**. A lancet **686** having a lancet tip **687** adjacent to the sample input port **682** is operably maintained within the housing such that the lancet **686** can move to extend the lancet tip **687** through the sample input port **682** to outside of the sampling module cartridge.

The lancet **686** also has a lancet head **688** opposite the lancet tip. The lancet **686** driven to move by a lancet driver **689**, which is schematically depicted as a coil around the lancet **686**. The lancet driver **689** optionally is included in the 10 sampling module cartridge as pictured or alternatively is external to the sampling module cartridge. The sampling module may further include a driver port **690** defined by the housing adjacent to the lancet head **688**—the driver port **690** allows an external lancet driver **691** access to the lancet **686**.

In embodiments where the lancet driver **689** is in the sampling module cartridge, it may be necessary to have a driver connection point **694** upon the housing accessible to the analyzer device. The driver connection point **694** may be a means of triggering the lancet driver **689** or of supplying motive 20 force to the lancet driver **689**, e.g. an electrical current to an electromechanical lancet driver. Note that any of the drivers discussed above, including controllable drivers, electromechanical drivers, etc., can be substituted for the lancet driver **689** shown.

In one embodiment a pierceable membrane 692 is present between the lancet tip 687 and the sample input port 682, sealing the lancet 686 from any outside contact prior to use. A second membrane 693 may be present adjacent to the lancet head 688 sealing the driver port 690. The pierceable membrane 692 and the second membrane 693 function to isolate the lancet 686 within the lancet chamber to maintain sterility of the lancet 686 prior to use. During use the lancet tip 687 and the external lancet driver 691 pierce the pierceable membrane 692 and the second membrane 693, if present respectively.

A sample flow channel 695 leads from the sample input port 682 to an analytical region 696. The analytical region 696 is associated with a sample analyte sensor capable of being read by the analyzer device. If the sample analyte sensor is optical in nature, the sample analyte sensor may include 40 optically transparent windows 697 in the housing above and below the analytical region 696, allowing a light source in the analyzer device to pass light 698 through the analytical region. An optical analyte sensor 698', e.g. a CMOS array, is present in the analyzer device for sensing the light 699 that 45 has passed through the analytical region 696 and generating a signal to be analyzed by the analyzer device.

In a separate embodiment, only one optically transparent window is present, and the opposing side of the analytical region is silvered or otherwise reflectively coated to reflect 50 light back through the analytical region and out the window to be analyzed by the analyzer device. In an alternate embodiment, the analyte sensor is electrochemical **700**, e.g. an enzyme electrode, and includes a means of transmitting an electric current from the sampling module cartridge to the 55 analyzer device, e.g. an electrical contact **701**, or plurality of electrical contacts **701**, on the housing accessible to the analyzer device.

In one embodiment, the pierceable membrane **692** may be made of polymer-based film that has been coated with a 60 silicone-based gel. For example, the membrane structure may comprise a polymer-based film composed of polyethylene terephthalate, such as the film sold under the trademark MYLAR®. The membrane structure may further comprise a thin coating of a silicone-based gel such as the gel sold under 65 the trademark SYLGARD® on at least one surface of the film.

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The usefulness of such a film is its ability to reseal after the lancet tip has penetrated it without physically affecting the lancet's cutting tip and edges. The MYLAR® film provides structural stability while the thin SYLGARD® silicone laminate is flexible enough to retain its form and close over the hole made in the MYLAR® film. Other similar materials fulfilling the structural stability and flexibility roles may be used in the manufacture of the pierceable membrane in this embodiment.

The pierceable membrane 692 operates to allow the lancet tip 687 to pierce the pierceable membrane 692 as the lancet tip 687 travels into and through the sampling port 682. In the described embodiment, the silicone-based gel of the membrane 692 automatically seals the cut caused by the lancet tip 687. Therefore, after an incision is made on a finger of a user and the lancet tip 687 is retracted back through the pierceable membrane 692, the blood from the incision is prevented from flowing through the pierceable membrane 692, which aids the blood to travel through the sample flow channel 695 to accumulate within the analytical region 696.

Thus the pierceable membrane 692 prevents blood from flowing into the lancet device assembly, and blood contamination and loss into the lancet device mechanism cavity are prevented. In yet another embodiment, used sample input ports are automatically sealed off before going to the next sample acquisition cycle by a simple button mechanism. A similar mechanism seals off a sample input port should sampling be unsuccessful.

In an alternate embodiment, a calibrant supply reservoir 702 is also present in each sampling module. The calibrant supply reservoir 702 is filled with a calibrant solution and is in fluid communication with a calibration chamber 703. The calibration chamber 703 provides a source of a known signal from the sampling module cartridge to be used to validate and quantify the test conducted in the analytical region 696. As such, the configuration of the calibration chamber 703 closely resembles the analytical region 696.

During use, the calibrant solution is forced from the calibrant supply reservoir 702 into the calibration chamber 703. The figure depicts a stylized plunger 704 above the calibrant supply reservoir 702 ready to squeeze the calibrant supply reservoir 702. In practice, a variety of methods of transporting small quantities of fluid are known in the art and can be implemented on the sampling module cartridge. The calibration chamber 703 is associated with a calibrant testing means.

FIG. 73 shows two alternate calibrant testing means—optical windows 697 and an electrochemical analyte sensor 676. In cases where the sampling module is designed to perform several different tests on the blood, both optical and electrochemical testing means may be present. The optical windows 697 allow passage of light 677 from the analyzer device through the calibration chamber 703, whereupon the light 703' leaving the calibration chamber 703 passes onto an optical analyte sensor 698' to result in a signal in the analyzer device.

The electrochemical analyte sensor 676 is capable of generating a signal that is communicated to the analyzer device via, e.g. an electrical contact 704', which is accessible to a contact probe 702' on the analyzer device that can be extended to contact the electrical contact 704'. The calibrant solution may be any solution, which, in combination with the calibrant testing means, will provide a suitable signal, which will serve as calibration measurement to the analyzer device. Suitable calibrant solutions are known in the art, e.g. glucose solutions of known concentration. The calibration measurement is used to adjust the results obtained from sample analyte sensor from the analytical region 696.

To maintain small size in some sampling module cartridge embodiments, allowing small quantities of sampled blood to be sufficient, each component of the sampling module must be small, particularly the sample flow channel and the analytical region. The sample flow channel can be less than about 5 0.5 mm in diameter, specifically less than about 0.3 mm in diameter, more specifically less than about 0.2 mm in diameter, and even more specifically less than about 0.1 mm in diameter.

The sample flow channel may generally be at least about 50 micrometers in diameter. The dimensions of the analytical region may be less than about 1 mm by about 1 mm by about 1 mm, specifically less than about 0.6 mm by about 0.4 mm, more specifically less than about 0.4 mm by 0.4 mm by 0.2 mm, and even more specifically less than about 0.5 mm by about 0.2 mm by about 0.1 mm. The analytical region can generally be at least about 100 micrometers by 100 micrometers by 50 micrometers.

The sampling module cartridge is able to return a valid testing result with less than about 5 microliters of blood taken 20 from the skin of a patient, specifically less than about 1 microliter, more specifically less than about 0.4 microliters, and even more specifically less than about 0.2 microliters. Generally, at least 0.05 microliters of blood is drawn for a sample.

The cartridge housing may be made in a plurality of distinct pieces, which are then assembled to provide the completed housing. The distinct pieces may be manufactured from a wide range of substrate materials. Suitable materials for forming the described apparatus include, but are not limited 30 to, polymeric materials, ceramics (including aluminum oxide and the like), glass, metals, composites, and laminates thereof. Polymeric materials are particularly preferred herein and will typically be organic polymers that are homopolymers or copolymers, naturally occurring or synthetic, cross 35 linked or uncross linked.

It is contemplated that the various components and devices described herein, such as sampling module cartridges, sampling modules, housings, etc., may be made from a variety of materials, including materials such as the following: polycar- 40 bonates; polyesters, including poly (ethylene terephthalate) and poly(butylene terephthalate); polyamides, (such as nylons); polyethers, including polyformaldehyde and poly (phenylene sulfide); polyimides, such as that manufactured under the trademarks KAPTON (DuPont, Wilmington, Del.) 45 and UPILEX (Ube Industries, Ltd., Japan); polyolefin compounds, including ABS polymers, Kel-F copolymers, poly (methyl methacrylate), poly(styrene-butadiene) copolymers, poly(tetrafluoroethylene), poly(ethylenevinyl acetate) copolymers, poly(N-vinylcarbazole) and polystyrene.

The various components and devices described herein may also be fabricated from a "composite," i.e., a composition comprised of unlike materials. The composite may be a block composite, e.g., an A-B-A block composite, an A-B-C block composite, or the like. Alternatively, the composite may be a 55 heterogeneous combination of materials, i.e., in which the materials are distinct from separate phases, or a homogeneous combination of unlike materials. A laminate composite with several different bonded layers of identical or different materials can also be used.

Other preferred composite substrates include polymer laminates, polymer-metal laminates, e.g., polymer coated with copper, a ceramic-in-metal or a polymer-in-metal composite. One composite material is a polyimide laminate formed from a first layer of polyimide such as KAPTON 65 polyimide, available from DuPont (Wilmington, Del.), that has been co-extruded with a second, thin layer of a thermal

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adhesive form of polyimide known as KJ®, also available from DuPont (Wilmington, Del.).

Any suitable fabrication method for the various components and devices described herein can be used, including, but not limited to, molding and casting techniques, embossing methods, surface machining techniques, bulk machining techniques, and stamping methods. Further, injection-molding techniques well known in the art may be useful in shaping the materials used to produce sample modules and other components.

For some embodiments, the first time a new sampling module cartridge 668 is used, the user removes any outer packaging material from the sampling module cartridge 668 and opens the lid 671 of the analyzer device 669, exposing the chamber. The sampling module cartridge 668 is slipped into the chamber and the lid 671 closed. The patient's skin is positioned upon the sampling site 678 and the integrated process of lancing the skin, collecting the blood sample, and testing the blood sample is initiated, e.g. by pressing a function button 674 to cause the lancet driver to be triggered. The patient's skin is maintained in position upon the sampling site 678, adjacent the sample input port 682, until an adequate volume of blood has been collected, whereupon the system may emit a signal (e.g. an audible beep) that the patient's skin may be lifted from the sampling site 678.

When the testing of the sample is complete, the analyzer device 669 automatically reads the results from the sampling module cartridge 668 and reports the results on the readout display 673. The analyzer device 669 may also store the result in memory for later downloading to a computer system. The sampling module cartridge 668 may then automatically be advanced to bring the next sampling module inline for the next use. Each successive time the system is used (optionally until the sampling module cartridge 668 is used up), the patient's skin may be placed upon the sampling site 678 of the (already installed) sampling module cartridge 668, thus simplifying the process of blood sampling and testing.

A method of providing more convenient blood sampling, wherein a series of blood samples may be collected and tested using a single disposable sampling module cartridge which is designed to couple to an analyzer device is described. Embodiments of the sampling module cartridge include a plurality of sampling modules. Each sampling module can be adapted to perform a single blood sampling cycle and is functionally arranged within the sampling module cartridge to allow a new sampling module to be brought online after a blood sampling cycle is completed.

Each blood sampling cycle may include lancing of a patient's skin, collection of a blood sample, and testing of the blood sample. The blood sampling cycle may also include reading of information about the blood sample by the analyzer device, display and/or storage of test results by the analyzer device, and/or automatically advancing the sampling module cartridge to bring a new sampling module 55 online and ready for the next blood sampling cycle to begin.

A method embodiment starts with coupling of the sampling module cartridge and analyzer device and then initiating a blood sampling cycle. Upon completion of the blood sampling cycle, the sampling module cartridge is advanced to bring a fresh, unused sampling module online, ready to perform another blood sampling cycle. Generally, at least ten sampling modules are present, allowing the sampling module cartridge to be advanced nine times after the initial blood sampling cycle.

In some embodiments, more sampling modules are present and the sampling module cartridge may be advanced about 19 times, and about 34 times in some embodiments, allowing

about 19 or about 34 blood sampling cycles, respectively, after the initial blood sampling cycle. After a series of blood sampling cycles has been performed and substantially all (i.e. more than about 80%) of the sampling modules have been used, the sampling module cartridge is decoupled from the analyzer device and discarded, leaving the analyzer device ready to be coupled with a new sampling module cartridge.

Referring to FIGS. 74-76, a tissue penetration sampling device 180 is shown with the controllable driver 179 of FIG. 20 coupled to a sampling module cartridge 705 and disposed within a driver housing 706. A ratchet drive mechanism 707 is secured to the driver housing 706, coupled to the sampling module cartridge 705 and configured to advance a sampling module belt 708 within the sampling module cartridge 705 so as to allow sequential use of each sampling module 709 in the sampling module belt 708. The ratchet drive mechanism 707 has a drive wheel 711 configured to engage the sampling modules 709 of the sampling module belt 708. The drive wheel 711 is coupled to an actuation lever 712 that advances the drive wheel 711 in increments of the width of a single 20 sampling module 709. A T-slot drive coupler 713 is secured to the elongated coupler shaft 184.

A sampling module **709** is loaded and ready for use with the drive head **198** of the lancet **183** of the sampling module **709** loaded in the T-slot **714** of the drive coupler **713**. A 25 sampling site **715** is disposed at the distal end **716** of the sampling module **709** disposed about a lancet exit port **717**. The distal end **716** of the sampling module **709** is exposed in a module window **718**, which is an opening in a cartridge cover **721** of the sampling module cartridge **705**. This allows the distal end **716** of the sampling module **709** loaded for use to be exposed to avoid contamination of the cartridge cover **721** with blood from the lancing process.

A reader module 722 is disposed over a distal portion of the sampling module 709 that is loaded in the drive coupler 713 35 for use and has two contact brushes 724 that are configured to align and make electrical contact with analyte sensor contacts 725 of the sampling module 709 as shown in FIG. 77. With electrical contact between the analyte sensor contacts 725 and contact brushes 724, the processor 193 of the controllable 40 driver 179 can read a signal from an analytical region 726 of the sampling module 709 after a lancing cycle is complete and a blood sample enters the analytical region 726 of the sampling module 709. The contact brushes 724 can have any suitable configuration that will allow the sampling module 45 belt 708 to pass laterally beneath the contact brushes 724 and reliably make electrical contact with the sampling module 709 loaded in the drive coupler 713 and ready for use. A spring loaded conductive ball bearing is one example of a contact brush 724 that could be used. A resilient conductive 50 strip shaped to press against the inside surface of the flexible polymer sheet 727 along the analyte sensor contact region 728 of the sampling module 709 is another embodiment of a contact brush 724.

The sampling module cartridge **705** has a supply canister **729** and a receptacle canister **730**. The unused sampling modules of the sampling module belt **708** are disposed within the supply canister **729** and the sampling modules of the sampling module belt **708** that have been used are advanced serially after use into the receptacle canister **730**.

FIG. 77 is a perspective view of a section of the sampling module belt 708 shown in the sampling module cartridge 705 in FIG. 74. The sampling module belt 708 has a plurality of sampling modules 709 connected in series by a sheet of flexible polymer 727. The sampling module belt 708 shown in 65 FIG. 77 is formed from a plurality of sampling module body portions 731 that are disposed laterally adjacent each other

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and connected and sealed by a single sheet of flexible polymer 727. The flexible polymer sheet 727 can optionally have analyte sensor contacts 725, flexible electrical conductors 732, sample analyte sensors 733 or any combination of these elements formed on the inside surface 734 of the flexible polymer sheet 727. These electrical, optical or chemical elements can be formed by a variety of methods including vapor deposition and the like.

The proximal portion 735 of the flexible polymer sheet 727 has been folded over on itself in order to expose the analyte sensor contacts 725 to the outside surface of the sampling module 709. This makes electrical contact between the contact brushes 724 of the reader module 722 and the analyte sensor contacts 725 easier to establish as the sampling modules 709 are advanced and loaded into position with the drive coupler 713 of the controllable driver 179 ready for use. The flexible polymer sheet 727 can be secured to the sampling module body portion 731 by adhesive bonding, solvent bonding, ultrasonic thermal bonding or any other suitable method.

FIG. **78** shows a perspective view of a single sampling module **709** of the sampling module belt **708** of FIG. **77** during the assembly phase of the sampling module **709**. The proximal portion **735** of the flexible polymer sheet **727** is being folded over on itself as shown in order to expose the analyte sensor contacts **725** on the inside surface of the flexible polymer sheet **727**. FIG. **79** is a bottom view of a section of the flexible polymer sheet **727** of the sampling module **709** of FIG. **78** illustrating the analyte sensor contacts **725**, flexible conductors **732** and sample analyte sensors **733** deposited on the bottom surface of the flexible polymer sheet **727**.

A lancet 183 is shown disposed within the lancet channel 736 of the sampling module 709 of FIG. 78 as well as within the lancet channels 736 of the sampling modules 709 of the sampling module belt 708 of FIG. 77. The lancet 183 has a tip 196 and a shaft portion 201 and a drive head 198. The shaft portion 201 of the lancet slides within the lancet channel 736 of the sampling module 709 and the drive head 198 of the lancet 183 has clearance to move in a proximal and distal direction within the drive head slot 737 of the sampling module 709. Disposed adjacent the drive head slot 737 and at least partially forming the drive head slot are a first protective strut 737' and a second protective strut 737" that are elongated and extend substantially parallel to the lancet 183.

In one lancet 183 embodiment, the drive head 198 of the lancet 183 can have a width of about 0.9 to about 1.1 mm. The thickness of the drive head 198 of the lancet 183 can be about 0.4 to about 0.6 mm. The drive head slot 714 of the sampling module 709 should have a width that allows the drive head 198 to move freely within the drive head slot 714. The shaft portion 201 of the lancet 183 can have a transverse dimension of about 50 µm to about 1000 µm. Typically, the shaft portion 201 of the lancet 183 has a round transverse cross section, however, other configurations are contemplated.

The sampling module body portions 731 and the sheet of flexible polymer 727 can both be made of polymethylmethacrylate (PMMA), or any other suitable polymer, such as those discussed above. The dimensions of a typical sampling module body portion 731 can be about 14 to about 18 mm in length, about 4 to about 5 mm in width, and about 1.5 to about 2.5 mm in thickness. In other embodiments, the length of the sample module body portion can be about 0.5 to about 2.0 inch and the transverse dimension can be about 0.1 to about 0.5 inch. The thickness of the flexible polymer sheet 727 can be about 100 to about 150 microns. The distance between adjacent sampling modules 709 in the sampling module belt 708 can vary from about 0.1 mm to about 0.3 mm, and in some embodiments, from about 0.2 to about 0.6.

FIGS. **80** and **81** show a perspective view of the body portion **731** of the sampling module **709** of FIG. **77** without the flexible polymer cover sheet **727** or lancet **183** shown for purposes of illustration. FIG. **81** is an enlarged view of a portion of the body portion **731** of the sampling module **709** of FIG. **80** illustrating the sampling site **715**, sample input cavity **715**', sample input port **741**, sample flow channel **742**, analytical region **743**, control chamber **744**, vent **762**, lancet channel **736**, lancet channel stopping structures **747** and **748** and lancet guides **749-751** of the sampling module **709**.

The lancet channel 736 has a proximal end 752 and a distal end 753 and includes a series of lancet bearing guide portions 749-751 and sample flow stopping structures 747-748. The lancet guides 749-751 may be configured to fit closely with the shaft of the lancet 183 and confine the lancet 183 to substantially axial movement. At the distal end 753 of the lancet channel 736 the distal-most lancet guide portion 749 is disposed adjacent the sample input port 741 and includes at its distal-most extremity, the lancet exit port 754 which is disposed adjacent the sample input cavity 715'. The sample input 20 cavity can have a transverse dimension, depth or both, of about 2 to 5 times the transverse dimension of the lancet 183, or about 0.2 to about 2 mm, specifically, about 0.4 to about 1.5 mm, and more specifically, about 0.5 to about 1.0 mm. The distal-most lancet guide 749 can have inner transverse dimen- 25 sions of about 300 to about 350 microns in width and about 300 to about 350 microns in depth. Proximal of the distalmost lancet guide portion 749 is a distal sample flow stop 747 that includes a chamber adjacent the distal-most lancet 749. The chamber has a transverse dimension that is significantly 30 larger than the transverse dimension of the distal-most lancet guide 749. The chamber can have a width of about 600 to about 800 microns, and a depth of about 400 to about 600 microns and a length of about 2000 to about 2200 microns. The rapid transition of transverse dimension and cross sec- 35 tional area between the distal-most lancet bearing guide 749 and the distal sample flow stop 747 interrupts the capillary action that draws a fluid sample through the sample input cavity 715' and into the lancet channel 736.

A center lancet bearing guide 750 is disposed proximal of 40 the distal lancet channel stop 747 and can have dimensions similar to those of the distal-most lancet bearing guide 749. Proximal of the center lancet guide 750 is a proximal lancet channel stop 748 with a chamber. The dimensions of the proximal lancet channel stop can be the same or similar to 45 those of the distal lancet channel stop 747. The proximal lancet channel stop 748 can have a width of about 600 to about 800 microns, and a depth of about 400 to about 600 microns and a length of about 2800 to about 3000 microns. Proximal of the proximal lancet channel stop 748 is a proximal lancet 50 guide 751. The proximal lancet guide 751 can dimensions similar to those of the other lancet guide 749 and 750 portions with inner transverse dimensions of about 300 to about 350 microns in width and about 300 to about 350 microns in depth. Typically, the transverse dimension of the lancet 55 guides 749-751 are about 10 percent larger than the transverse dimension of the shaft portion 201 of the lancet 183 that the lancet guides 749-751 are configured to guide.

A proximal fracturable seal (not shown) can be positioned between the proximal lancet guide **751** and the shaft portion 60 **201** of the lancet **183** that seals the chamber of the proximal lancet channel stop **748** from the outside environment. The fracturable seal seals the chamber of the proximal lancet channel stop **748** and other interior portions of the sample chamber from the outside environment when the sampling 65 module **709** is stored for use. The fracturable seal remains intact until the lancet **183** is driven distally during a lancet

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cycle at which point the seal is broken and the sterile interior portion of the sample chamber is exposed and ready to accept input of a liquid sample, such as a sample of blood. A distal fracturable seal (not shown) can be disposed between the lancet 183 and the distal-most lancet guide 749 of the sampling module 709 to seal the distal end 753 of the lancet channel 736 and sample input port 741 to maintain sterility of the interior portion of the sampling module 709 until the lancet 183 is driven forward during a lancing cycle.

Adjacent the lancet exit port 754 within the sample input cavity 715' is the sample input port 741 that is configured to accept a fluid sample that emanates into the sample input cavity 715' from target tissue 233 at a lancing site after a lancing cycle. The dimensions of the sample input port 741 can a depth of about 60 to about 70 microns, a width of about 400 to about 600 microns. The sample input cavity can have a transverse dimension of about 2 to about 5 times the transverse dimension of the lancet 183, or about 400 to about 1000 microns. The sample input cavity serves to accept a fluid sample as it emanates from lanced tissue and direct the fluid sample to the sample input port 741 and thereafter the sample flow channel 742. The sample flow channel 742 is disposed between and in fluid communication with the sample input port 741 and the analytical region 743. The transverse dimensions of the sample flow channel 742 can be the same as the transverse dimensions of the sample input port 741 with a depth of about 60 to about 70 microns, a width of about 400 to about 600 microns. The length of the sample flow channel 742 can be about 900 to about 1100 microns. Thus, in use, target tissue is disposed on the sampling site 715 and a lancing cycle initiated. Once the target tissue 233 has been lanced and the sample begins to flow therefrom, the sample enters the sample input cavity 715' and then the sample input port 741. The sample input cavity 715' may be sized and configured to facilitate sampling success by applying pressure to a perimeter of target tissue 233 before, during and after the lancing cycle and hold the wound track open after the lancing cycle to allow blood or other fluid to flow from the wound track and into the sample input cavity 715'. From the sample input port 741, the sample in then drawn by capillary or other forces through the sample flow channel 742 and into the analytical region 743 and ultimately into the control chamber 744. The control chamber 744 may be used to provide indirect confirmation of a complete fill of the analytical region 743 by a sample fluid. If a fluid sample has been detected in the control chamber 744, this confirms that the sample has completely filled the analytical region 743. Thus, sample detectors may be positioned within the control chamber 744 to confirm filling of the analytical region 743.

The analytical region **743** is disposed between and in fluid communication with the sample flow channel **742** and the control chamber **744**. The analytical region **743** can have a depth of about 60 to about 70 microns, a width of about 900 to about 1100 microns and a length of about 5 to about 6 mm. A typical volume for the analytical region **743** can be about 380 to about 400 nanoliters. The control chamber **744** is disposed adjacent to and proximal of the analytical region **743** and can have a transverse dimension or diameter of about 900 to about 1100 microns and a depth of about 60 to about 70 microns.

The control chamber **744** is vented to the chamber of the proximal lancet channel stop **748** by a vent that is disposed between and in fluid communication with the control chamber **744** and the chamber of the proximal lancet channel stop **748**. Vent **762** can have transverse dimensions that are the same or similar to those of the sample flow channel **742** disposed between the analytical region **743** and the sample

input port **741**. Any of the interior surfaces of the sample input port **741**, sample flow channels **742** and **762**, analytical region **743**, vents **745** or control chamber **744** can be coated with a coating that promotes capillary action. A hydrophilic coating such as a detergent is an example of such a coating.

The analytical region **743** accommodates a blood sample that travels by capillary action from the sampling site **715** through the sample input cavity **715**' and into the sample input port **741**, through the sample flow channel **742** and into the analytical region **743**. The blood can then travel into the control chamber **744**. The control chamber **744** and analytical region **743** are both vented by the vent **762** that allows gases to escape and prevents bubble formation and entrapment of a sample in the analytical region **743** and control chamber **744**. Note that, in addition to capillary action, flow of a blood sample into the analytical region **743** can be facilitated or accomplished by application of vacuum, mechanical pumping or any other suitable method.

Once a blood sample is disposed within the analytical region **743**, analytical testing can be performed on the sample 20 with the results transmitted to the processor **193** by electrical conductors **732**, optically or by any other suitable method or means. In some embodiments, it may be desirable to confirm that the blood sample has filled the analytical region **743** and that an appropriate amount of sample is present in the chamber in order to carry out the analysis on the sample.

Confirmation of sample arrival in either the analytical region 743 or the control chamber 744 can be achieved visually, through the flexible polymer sheet 727 which can be transparent. However, it may be desirable in some embodi- 30 ments to use a very small amount of blood sample in order to reduce the pain and discomfort to the patient during the lancing cycle. For sampling module 709 embodiments such as described here, having the sample input cavity 715' and sample input port 741 adjacent the lancet exit port 754 allows 35 the blood sample to be collected from the patient's skin 233 without the need for moving the sampling module 709 between the lancing cycle and the sample collection process. As such, the user does not need to be able to see the sample in Because of this, the position of the sample input cavity 715' and the sample input port 741 adjacent the lancet exit port 754 allows a very small amount of sample to be reliably obtained and tested.

Samples on the order of tens of nanoliters, such as about 10 to about 50 nanoliters can be reliably collected and tested with a sampling module **709**. This size of blood sample is too small to see and reliably verify visually. Therefore, it is necessary to have another method to confirm the presence of the blood sample in the analytical region **743**. Sample analyte sensors 50 **733**, such as the thermal sample analyte sensors discussed above can positioned in the analytical region **743** or control chamber **744** to confirm the arrival of an appropriate amount of blood sample.

In addition, optical methods, such as spectroscopic analysis of the contents of the analytical region **743** or control chamber **744** could be used to confirm arrival of the blood sample. Other methods such as electrical detection could also be used and these same detection methods can also be disposed anywhere along the sample flow path through the sampling module **709** to confirm the position or progress of the sample (or samples) as it moves along the flow path as indicated by the arrows **763** in FIG. **81**. The detection methods described above can also be useful for analytical methods requiring an accurate start time.

The requirement for having an accurate start time for an analytical method can in turn require rapid filling of an ana-

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lytical region 743 because many analytical processes begin once the blood sample enters the analytical region 743. If the analytical region 743 takes too long to fill, the portion of the blood sample that first enters the analytical region 743 will have been tested for a longer time that the last portion of the sample to enter the analytical region 743 which can result in inaccurate results. Therefore, it may be desirable in these circumstances to have the blood sample flow first to a reservoir, filling the reservoir, and then have the sample rapidly flow all at once from the reservoir into the analytical region 743.

In one embodiment of the sampling module 709, the analytical region 743 can have a transverse cross section that is substantially greater than a transverse cross section of the control chamber 744. The change in transverse cross section can be accomplished by restrictions in the lateral transverse dimension of the control chamber 744 versus the analytical region 743, by step decreases in the depth of the control chamber 744, or any other suitable method. Such a step between the analytical region 743 and the control chamber 744 is shown in FIG. 81. In such an embodiment, the analytical region 743 can behave as a sample reservoir and the control chamber 744 as an analytical region that requires rapid or nearly instantaneous filling in order to have a consistent analysis start time. The analytical region 743 fills by a flow of sample from the sample flow channel 742 until the analytical region is full and the sample reaches the step decrease in chamber depth at the boundary with the control chamber 744. Once the sample reaches the step decrease in cross sectional area of the control chamber 744, the sample then rapidly fills the control chamber 744 by virtue of the enhanced capillary action of the reduced cross sectional area of the control chamber 744. The rapid filling of the control chamber allows any analytical process initiated by the presence of sample to be carried out in the control chamber 744 with a reliable start time for the analytical process for the entire sample of the control chamber 744.

Filling by capillary force is passive. It can also be useful for some types of analytical testing to discard the first portion of a sample that enters the sampling module 709, such as the case where there may be interstitial fluid contamination of the sample input port 741 adjacent the lancet exit port 754 allows a very small amount of sample to be reliably obtained and tested.

Samples on the order of tens of nanoliters, such as about 10 to about 50 nanoliters can be reliably collected and tested with a sampling module 709. This size of blood sample is too small to see and reliably verify visually. Therefore, it is necessary to have another method to confirm the presence of the blood

For some types of analytical testing, it may be advantageous to have multiple analytical regions **743** in a single sampling module **709**. In this way multiple iterations of the same type of analysis could be performed in order to derive some statistical information, e.g. averages, variation or confirmation of a given test or multiple tests measuring various different parameters could be performed in different analytical regions **743** in the same sampling module **709** filled with a blood sample from a single lancing cycle.

FIG. 82 is an enlarged elevational view of a portion of an alternative embodiment of a sampling module 766 having a plurality of small volume analytical regions 767. The small volume analytical regions 767 can have dimensions of about 40 to about 60 microns in width in both directions and a depth that yields a volume for each analytical region 767 of about 1 nanoliter to about 100 nanoliters, specifically about 10 nanoliters to about 50 nanoliters. The array of small volume analytical regions 767 can be filled by capillary action through a

sample flow channel **768** that branches at a first branch point **769**, a second branch point **770** and a third branch point **771**. Each small volume analytical region **767** can be used to perform a like analytical test or a variety of different tests can be performed in the various analytical regions **767**.

For some analytical tests, the analytical regions 767 must have maintain a very accurate volume, as some of the analytical tests that can be performed on a blood sample are volume dependent. Some analytical testing methods detect glucose levels by measuring the rate or kinetic of glucose 10 consumption. Blood volume required for these tests is on the order of about 1 to about 3 microliters. The kinetic analysis is not sensitive to variations in the volume of the blood sample as it depends on the concentration of glucose in the relatively large volume sample with the concentration of glucose 15 remaining essentially constant throughout the analysis. Because this type of analysis dynamically consumes glucose during the testing, it is not suitable for use with small samples, e.g. samples on the order of tens of nanoliters where the consumption of glucose would alter the concentration of glu-

Another analytical method uses coulomb metric measurement of glucose concentration. This method is accurate if the sample volume is less than about 1 microliter and the volume of the analytical region is precisely controlled. The accuracy 25 and the speed of the method is dependent on the small and precisely known volume of the analytical region **767** because the rate of the analysis is volume dependent and large volumes slow the reaction time and negatively impact the accuracy of the measurement.

Another analytical method uses an optical fluorescence decay measurement that allows very small sample volumes to be analyzed. This method also requires that the volume of the analytical region 767 be precisely controlled. The small volume analytical regions 767 discussed above can meet the 35 criteria of maintaining small accurately controlled volumes when the small volume analytical regions 767 are formed using precision manufacturing techniques. Accurately formed small volume analytical regions 767 can be formed in materials such as PMMA by methods such as molding and 40 stamping. Machining and etching, either by chemical or laser processes can also be used. Vapor deposition and lithography can also be used to achieve the desired results.

The sampling modules **709** and **766** discussed above all are directed to embodiments that both house the lancet **183** and 45 have the ability to collect and analyze a sample. In some embodiments of a sampling module, the lancet **183** may be housed and a sample collected in a sample reservoir without any analytical function. In such an embodiment, the analysis of the sample in the sample reservoir may be carried out by 50 transferring the sample from the reservoir to a separate analyzer. In addition, some modules only serve to house a lancet **183** without any sample acquisition capability at all. The body portion **774** of such a lancet module **775** is shown in FIG. **83**. The lancet module **775** has an outer structure similar 55 to that of the sampling modules **709** and **766** discussed above, and can be made from the same or similar materials.

A flexible polymer sheet 727 (not shown) can be used to cover the face of the lancet module 775 and contain the lancet 183 in a lancet channel 776 that extends longitudinally in the 60 lancet module body portion 774. The flexible sheet of polymer 727 can be from the same material and have the same dimensions as the flexible polymer sheet 727 discussed above. Note that the proximal portion of the flexible polymer sheet 727 need not be folded over on itself because there are 65 no analyte sensor contacts 725 to expose. The flexible polymer sheet 727 in such a lancet module 775 serves only to

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confine the lancet 183 in the lancet channel 776. The lancet module 775 can be configured in a lancet module belt, similar to the sampling module belt 708 discussed above with the flexible polymer sheet 727 acting as the belt. A drive head slot 777 is dispose proximal of the lancet channel 776.

With regard to the tissue penetration sampling device 180 of FIG. 74, use of the device 180 begins with the loading of a sampling module cartridge 705 into the controllable driver housing 706 so as to couple the cartridge 705 to the controllable driver housing 706 and engage the sampling module belt 708 with the ratchet drive 707 and drive coupler 713 of the controllable driver 179. The drive coupler 713 can have a T-slot configuration such as shown in FIGS. 84 and 85. The distal end of the elongate coupler shaft 184 is secured to the drive coupler 713 which has a main body portion 779, a first and second guide ramp 780 and 781 and a T-slot 714 disposed within the main body portion 779. The T-slot 714 is configured to accept the drive head 198 of the lancet 183. After the sampling module cartridge 705 is loaded into the controllable driver housing 706, the sampling module belt 708 is advanced laterally until the drive head 198 of a lancet 183 of one of the sampling modules 709 is fed into the drive coupler 713 as shown in FIGS. 86-88. FIGS. 86-88 also illustrate a lancet crimp device 783 that bends the shaft portion 201 of a used lancet 183 that is adjacent to the drive coupler 713. This prevents the used lancet 183 from moving out through the module body 731 and being reused.

As the sampling modules 709 of the sampling module belt 708 are used sequentially, they are advanced laterally one at a time into the receptacle canister 730 where they are stored until the entire sampling module belt 708 is consumed. The receptacle canister 730 can then be properly disposed of in accordance with proper techniques for disposal of blood-contaminated waste. The sampling module cartridge 705 allows the user to perform multiple testing operations conveniently without being unnecessarily exposed to blood waste products and need only dispose of one cartridge after many uses instead of having to dispose of a contaminated lancet 183 or module 709 after each use.

FIGS. **89** and **90** illustrate alternative embodiments of sampling module cartridges. FIG. **89** shows a sampling module cartridge **784** in a carousel configuration with adjacent sampling modules **785** connected rigidly and with analyte sensor contacts **786** from the analytical regions of the various sampling modules **785** disposed near an inner radius **787** of the carousel. The sampling modules **785** of the sampling module cartridge **784** are advanced through a drive coupler **713** but in a circular as opposed to a linear fashion.

FIG. 90 illustrates a block of sampling modules 788 in a four by eight matrix. The drive head 198 of the lancets 183 of the sampling modules 789 shown in FIG. 90 are engaged and driven using a different method from that of the drive coupler 713 discussed above. The drive heads 198 of the lancets 183 have an adhesive coating 790 that mates with and secures to the drive coupler 791 of the lancet driver 179, which can be any of the drivers, including controllable drivers, discussed above.

The distal end 792 of the drive coupler 791 contacts and sticks to the adhesive 790 of proximal surface of the drive head 198 of the lancet 183 during the beginning of the lancet cycle. The driver coupler 791 pushes the lancet 183 into the target tissue 237 to a desired depth of penetration and stops. The driver coupler 791 then retracts the lancet 183 from the tissue 233 using the adhesive contact between the proximal surface of the drive head 198 of the lancet 183 and distal end surface of the drive coupler 791, which is shaped to mate with the proximal surface.

At the top of the retraction stroke, a pair of hooked members 793 which are secured to the sampling module 789 engage the proximal surface of the drive head 198 and prevent any further retrograde motion by the drive head 198 and lancet 183. As a result, the drive coupler 791 breaks the 5 adhesive bond with the drive head 198 and can then be advanced by an indexing operation to the next sampling module 789 to be used.

FIG. 91 is a side view of an alternative embodiment of a drive coupler 796 having a lateral slot 797 configured to 10 accept the L-shaped drive head 798 of the lancet 799 that is disposed within a lancet module 800 and shown with the L-shaped drive head 798 loaded in the lateral slot 797. FIG. 92 is an exploded view of the drive coupler 796, lancet 799 with L-shaped drive head 798 and lancet module 800 of FIG. 91. 15 This type of drive coupler 796 and drive head 798 arrangements could be substituted for the configuration discussed above with regard to FIGS. 84-88. The L-shaped embodiment of the drive head 798 may be a less expensive option for producing a coupling arrangement that allows serial advance- 20 ment of a sampling module belt or lancet module belt through the drive coupler **796** of a lancet driver, such as a controllable lancet driver 179.

For some embodiments of multiple lancing devices 180, it does not require a lancet module 775 in order to house the lancets 183 stored in a cartridge. Eliminating the lancet modules 775 from a multiple lancet device 180 allows for a higher capacity cartridge because the volume of the cartridge is not taken up with the bulk of multiple modules 775. FIGS. 93-96 30 illustrate a high capacity lancet cartridge coupled to a belt advance mechanism 804. The belt advance mechanism 804 is secured to a controlled driver 179 housing which contains a controlled electromagnetic driver.

The lancet cartridge 803 has a supply canister 805 and a 35 receptacle canister 806. A lancet belt 807 is disposed within the supply canister 805. The lancet belt 807 contains multiple sterile lancets 183 with the shaft portion 201 of the lancets 183 disposed between the adhesive surface 808 of a first carrier tape 809 and the adhesive surface 810 of a second 40 carrier tape 811 with the adhesive surfaces 808 and 810 pressed together-around the shaft portion 201 of the lancets 183 to hold them securely in the lancet belt 807. The lancets 183 have drive heads 198 which are configured to be laterally engaged with a drive coupler 713, which is secured to an 45 elongate coupler shaft 184 of the controllable driver 179.

The belt advance mechanism 804 includes a first cog roller 814 and a second cog roller 815 that have synchronized rotational motion and are advanced in unison in an incremental indexed motion. The indexed motion of the first and sec- 50 ond cog rollers 814 and 815 advances the lancet belt 807 in units of distance equal to the distance between the lancets 183 disposed in the lancet belt 807. The belt advance mechanism 804 also includes a first take-up roller 816 and a second take-up roller 817 that are configured to take up slack in the 55 first and second carrier tapes 809 and 811 respectively.

When a lancet belt cartridge 803 is loaded in the belt advance mechanism 804, a lead portion 818 of the first carrier tape 809 is disposed between a first cog roller 814 and a second cog roller 815 of the belt advance mechanism 804. 60 The lead portion 818 of the first carrier tape 809 wraps around the outer surface 819 of the first turning roller 827, and again engages roller 814 with the cogs 820 of the first cog roller 814 engaged with mating holes 821 in the first carrier tape 809. The lead portion 818 of the first carrier tape 809 is then 65 secured to a first take-up roller 816. A lead portion 822 of the second carrier tape 811 is also disposed between the first cog

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roller 814 and second cog roller 815 and is wrapped around an outer surface 823 of the second turning roller 828, and again engages roller 815 with the cogs 826' of the second cog roller 815 engaged in with mating holes 825 of the second carrier tape 811. The lead portion 822 of the second carrier tape 811 is thereafter secured to a second take-up roller 817.

As the first and second cog rollers 814 and 815 are advanced, the turning rollers 827 and 828 peel the first and second carrier tapes 809 and 811 apart and expose a lancet 183. The added length or slack of the portions of the first and second carrier tapes 809 and 811 produced from the advancement of the first and second cog rollers 814 and 815 is taken up by the first and second take-up rollers 816 and 817. As a lancet 183 is peeled out of the first and second carrier tapes 809 and 811, the exposed lancet 183 is captured by a lancet guide wheel 826' of the belt advance mechanism 804, shown in FIG. 96, which is synchronized with the first and second cog rollers 814 and 815. The lancet guide wheel 826' then advances the lancet 183 laterally until the drive head 198 of the lancet 183 is loaded into the drive coupler 713 of the controllable driver 179. The controllable driver 179 can then be activated driving the lancet 183 into the target tissue 233 and retracted to complete the lancing cycle.

Once the lancing cycle is complete, the belt advance may be desirable to have a high capacity-lancing device that 25 mechanism 804 can once again be activated which rotates the lancet guide wheel 826 and advances the used lancet 183 laterally and into the receptacle canister 806. At the same time, a new unused lancet 183 is loaded into the drive coupler 713 and readied for the next lancing cycle. This repeating sequential use of the multiple lancing device 180 continues until all lancets 183 in the lancet belt 807 have been used and disposed of in the receptacle canister 806. After the last lancet 183 has been consumed, the lancet belt cartridge 803 can then be removed and disposed of without exposing the user to any blood contaminated materials. The belt advance mechanism 804 can be activated by a variety of methods, including a motorized drive or a manually operated thumbwheel which is coupled to the first and second cog rollers 814 and 815 and lancet guide wheel 826.

> Although discussion of the devices described herein has been directed primarily to substantially painless methods and devices for access to capillary blood of a patient, there are many other uses for the devices and methods. For example, the tissue penetration devices discussed herein could be used for substantially painless delivery of small amounts of drugs, or other bioactive agents such as gene therapy agents, vectors, radioactive sources etc. As such, it is contemplated that the tissue penetration devices and lancet devices discussed herein could be used to delivery agents to positions within a patient's body as well as taking materials from a patient's body such as blood, lymph fluid, spinal fluid and the like. Drugs delivered may include analgesics that would further reduce the pain perceived by the patient upon penetration of the patient's body tissue, as well as anticoagulants that may facilitate the successful acquisition of a blood sample upon penetration of the patient's tissue.

> Referring to FIGS. 97-101, a device for injecting a drug or other useful material into the tissue of a patient is illustrated. The ability to localize an injection or vaccine to a specific site within a tissue, layers of tissue or organ within the body can be important. For example, epithelial tumors can be treated by injection of antigens, cytokine, or colony stimulating factor by hypodermic needle or high-pressure injection sufficient for the antigen to enter at least the epidermis or the dermis of a patient. Often, the efficacy of a drug or combination drug therapy depends on targeted delivery to localized areas thus affecting treatment outcome.

The ability to accurately deliver drugs or vaccinations to a specific depth within the skin or tissue layer may avoid wastage of expensive drug therapies therefore impacting cost effectiveness of a particular treatment. In addition, the ability to deliver a drug or other agent to a precise depth can be a clear advantage where the outcome of treatment depends on precise localized drug delivery (such as with the treatment of intralesional immunotherapy). Also, rapid insertion velocity of a hypodermic needle to a precise predetermined depth in a patient's skin is expected to reduce pain of insertion of the needle into the skin. Rapid insertion and penetration depth of a hypodermic needle, or any other suitable elongated delivery device suitable for penetrating tissue, can be accurately controlled by virtue of a position feedback loop of a controllable driver coupled to the hypodermic needle.

FIG. 97 illustrates 901 distal end 901 of a hypodermic needle 902 being driven into layers of skin tissue 903 by an electromagnetic controllable driver 904. The electromagnetic controllable driver 904 of FIG. 79 can have any suitable configuration, such as the configuration of electromagnetic 20 controllable drivers discussed above. The layers of skin 903 being penetrated include the stratum corneum 905, the stratum lucidum 906, the stratum granulosum 907, the stratum spinosum 908, the stratum basale 909 and the dermis 911. The thickness of the stratum corneum 905 is typically about 300 25 micrometers in thickness. The portion of the epidermis excluding the stratum corneum 905 includes the stratum lucidum 906, stratum granulosum 907, and stratum basale can be about 200 micrometers in thickness. The dermis can be about 1000 micrometers in thickness. In FIG. 97, an outlet port 912 30 of the hypodermic needle 902 is shown disposed approximately in the stratum spinosum 908 layer of the skin 903 injecting an agent 913 into the stratum spinosum 908.

FIGS. 98-101 illustrate an agent injection module 915 including an injection member 916, that includes a collaps- 35 collapsible canister 917. ible canister 917 and the hypodermic needle 902, that may be driven or actuated by a controllable driver, such as any of the controllable drivers discussed above, to drive the hypodermic needle into the skin 903 for injection of drugs, vaccines or the like. The agent injection module 915 has a reservoir, which 40 can be in the form of the collapsible canister 917 having a main chamber 918, such as shown in FIG. 98, for the drug or vaccine 913 to be injected. A cassette of a plurality of agent injection modules 915 (not shown) may provide a series of metered doses for long-term medication needs. Such a cas- 45 sette may be configured similarly to the module cassettes discussed above. Agent injection modules 915 and needles 902 may be disposable, avoiding biohazard concerns from unspent drug or used hypodermic needles 902. The geometry of the cutting facets 921 of the hypodermic needle shown in 50 FIG. 79, may be the same or similar to the geometry of the cutting facets of the lancet 183 discussed above.

Inherent in the position and velocity control system of some embodiments of a controllable driver is the ability to precisely determine the position or penetration depth of the 55 hypodermic needle 902 relative to the controllable driver or layers of target tissue or skin 903 being penetrated. For embodiments of controllable drivers that use optical encoders for position analyte sensors, such as an Agilent HEDS 9200 series, and using a four edge detection algorithm, it is possible 60 to achieve an in plane spatial resolution of +/-17 µm in depth. If a total tissue penetration stroke is about 3 mm in length, such as might be used for intradermal or subcutaneous injection, a total of 88 position points can be resolved along the penetration stroke. A spatial resolution this fine allows precise placement of a distal tip 901 or outlet port 912 of the hypodermic needle 902 with respect to the layers of the skin

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903 during delivery of the agent or drug 913. In some embodiments, a displacement accuracy of better than about 200 microns can be achieved, in others a displacement accuracy of better than about 40 microns can be achieved.

The agent injection module 915 includes the injection member 916 which includes the hypodermic needle 902 and drug reservoir or collapsible canister 917, which may couple to an elongated coupler shaft 184 via a drive coupler 185 as shown. The hypodermic needle 902 can be driven to a desired penetration depth, and then the drug or other agent 913, such as a vaccine, is passed into an inlet port 922 of the needle 902 through a central lumen 923 of the hypodermic needle 902 as shown by arrow 924, shown in FIG. 98, and out of the outlet port 912 at the distal end 901 of the hypodermic needle 902, shown in FIG. 97.

Drug or agent delivery can occur at the point of maximum penetration, or following retraction of the hypodermic needle 902. In some embodiments, it may be desirable to deliver the drug or agent 913 during insertion of the hypodermic needle 902. Drug or agent delivery can continue as the hypodermic needle 902 is being withdrawn (this is commonly the practice during anesthesia in dental work). Alternatively drug delivery can occur while the needle 902 is stationary during any part of the retraction phase.

The hollow hypodermic needle 902 is fitted with the collapsible canister 917 containing a drug or other agent 913 to be dispensed. The walls 928 of this collapsible canister 917 can be made of a soft resilient material such as plastic, rubber, or any other suitable material. A distal plate 925 is disposed at the distal end 926 of the collapsible canister is fixed securely to the shaft 927 of the hypodermic needle proximal of the distal tip 901 of the hypodermic needle 902. The distal plate 925 is sealed and secured to the shaft 927 of the hypodermic needle 902 to prevent leakage of the medication 913 from the collapsible canister 917.

A proximal plate 931 disposed at a proximal end 932 of the collapsible canister 917 is slidingly fitted to a proximal portion 933 of the shaft 927 of the hypodermic needle 902 with a sliding seal 934. The sliding seal 934 prevents leakage of the agent or medication 913 between the seal 934 and an outside surface of the shaft 927 of the hypodermic needle 902. The sliding seal allows the proximal plate 931 of the collapsible canister 917 to slide axially along the needle 902 relative to the distal plate 925 of the collapsible canister 917. A drug dose may be loaded into the main chamber 918 of the collapsible canister 917 during manufacture, and the entire assembly protected during shipping and storage by packaging and guide fins 935 surrounding the drive head slot 936 of the agent injection module 915.

An injection cycle may begin when the agent injection module 915 is loaded into a ratchet advance mechanism (not shown), and registered at a drive position with a drive head 937 of the hypodermic needle 902 engaged in the drive coupler 185. The position of the hypodermic needle 902 and collapsible canister 917 in this ready position is shown in FIG. 99.

Once the drive head 937 of the agent injection module 915 is loaded into the driver coupler 185, the controllable driver can then be used to launch the injection member 916 including the hypodermic needle 902 and collapsible canister 917 towards and into the patient's tissue 903 at a high velocity to a pre-determined depth into the patients skin or other organ. The velocity of the injection member 916 at the point of contact with the patient's skin 903 or other tissue can be up to about 10 meters per second for some embodiments, specifically, about 2 to about 5 m/s. In some embodiments, the velocity of the injection-member 916 may be about 2 to about

10 m/s at the point of contact with the patient's skin 903. As the collapsible canister 917 moves with the hypodermic needle 902, the proximal plate 931 of the collapsible canister 917 passes between two latch springs 938 of module body 939 that snap in behind the proximal plate 931 when the 5 collapsible canister 917 reaches the end of the penetration stroke, as shown in FIG. 100.

The controllable driver then reverses, applies force in the opposite retrograde direction and begins to slowly (relative to the velocity of the penetration stroke) retract the hypodermic needle 902. The hypodermic needle 902 slides through the sliding seal 934 of the collapsible canister 917 while carrying the distal plate 925 of the collapsible canister with it in a proximal direction relative to the proximal plate 931 of the collapsible canister 917. This relative motion between the 15 distal plate 925 of the collapsible canister 917 and the proximal plate 931 of the collapsible canister 917 causes the volume of the main chamber 918 to decrease. The decreasing volume of the main chamber 918 forces the drug or other agent 913 disposed within the main chamber 918 of the col- 20 lapsible canister 917 out of the main chamber 918 into the inlet port 922 in the shaft 927 of the hypodermic needle 902. The inlet port 922 of the hypodermic needle 902 is disposed within an in fluid communication with the main chamber 918 of the collapsible canister 917 as shown in FIG. 80. The drug 25 or agent then passes through the central lumen 923 of the hollow shaft 927 of the hypodermic needle 902 and is then dispensed from the output port 912 at the distal end 901 of the hypodermic needle 902 into the target tissue 903. The rate of perfusion of the drug or other agent 913 may be determined 30 by an inside diameter or transverse dimension of the collapsible canister 917. The rate of perfusion may also be determined by the viscosity of the drug or agent 913 being delivered, the transverse dimension or diameter of the central lumen 923, the input port 922, or the output port 912 of the 35 hypodermic needle 902, as well as other parameters.

During the proximal retrograde retraction stroke of the hypodermic needle 902, drug delivery continues until the main chamber 918 of the collapsible canister 917 is fully collapsed as shown in FIG. 101. At this point, the drive 40 coupler 185 may continue to be retracted until the drive head 937 of the hypodermic needle 902 breaks free or the distal seal 941 between the distal plate 925 of the chamber and the hypodermic needle 902 fails, allowing the drive coupler 185 to return to a starting position. The distal tip 901 of the 45 hypodermic needle 902 can be driven to a precise penetration depth within the tissue 903 of the patient using any of the methods or devices discussed above with regard to achieving a desired penetration depth using a controllable driver or any other suitable driver.

In another embodiment, the agent injection module 915 is loaded into a ratchet advance mechanism that includes an adjustable or movable distal stage or surface (not shown) that positions the agent injection 915 module relative to a skin contact point or surface 942. In this way, an agent delivery 55 module 915 having a penetration stroke of predetermined fixed length, such as shown in FIGS. 99-101, reaches a presettable penetration depth. The movable stage remains stationary during a drug delivery cycle. In a variation of this embodiment, the moveable stage motion may be coordinated 60 with a withdrawal of the hypodermic needle 902 to further control the depth of drug delivery.

In another embodiment, the latch springs 938 shown in the agent injection module 915 of FIGS. 99-101 may be molded with a number of ratchet teeth (not shown) that engage the 65 proximal end 932 of the collapsible canister 917 as it passes by on the penetration stroke. If the predetermined depth of

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penetration is less than the full stroke, the intermediate teeth retain the proximal end 932 of the collapsible canister 917 during the withdrawal stroke in order to collapse the main chamber 918 of the collapsible canister 917 and dispense the drug or agent 913 as discussed above.

In yet another embodiment, drive fingers (not shown) are secured to an actuation mechanism (not shown) and replace the latch springs 938. The actuation mechanism is driven electronically in conjunction with the controllable driver by a processor or controller, such as the processor 60 discussed above, to control the rate and amount of drug delivered anywhere in the actuation cycle. This embodiment allows the delivery of medication during the actuation cycle as well as the retraction cycle.

Inherent in the position and velocity control system of a controllable driver is the ability to precisely define the position in space of the hypodermic needle 902, allowing finite placement of the hypodermic needle in the skin 903 for injection of drugs, vaccines or the like. Drug delivery can be discrete or continuous depending on the need.

FIGS. 102-106 illustrate an embodiment of a cartridge 945 that may be used for sampling that has both a lancet cartridge body 946 and an sampling cartridge body 947. The sampling cartridge body 947 includes a plurality of sampling module portions 948 that are disposed radially from a longitudinal axis 949 of the sampling cartridge body 947. The lancet cartridge body 946 includes a plurality of lancet module portions 950 that have a lancet channel 951 with a lancet 183 slidably disposed therein. The lancet module portions 950 are disposed radially from a longitudinal axis 952 of the lancet cartridge body 946.

The sampling cartridge body 947 and lancet cartridge body 946 are disposed adjacent each other in an operative configuration such that each lancet module portion 950 can be readily aligned in a functional arrangement with each sampling module portion 948. In the embodiment shown in FIGS. 102-106, the sampling cartridge body 947 is rotatable with respect to the lancet cartridge body 946 in order to align any lancet channel 951 and corresponding lancet 183 of the lancet cartridge body 946 with any of the lancet channels 953 of the sampling module portions 948 of the sampling cartridge body 947. The operative configuration of the relative location and rotatable coupling of the sampling cartridge body 947 and lancet cartridge body 946 allow ready alignment of lancet channels 951 and 953 in order to achieve a functional arrangement of a particular lancet module portion 950 and sampling module portion 948. For the embodiment shown, the relative motion used to align the particular lancet module portions 950 and sampling module portions 948 is confined to a single degree of freedom via relative rotation.

The ability of the cartridge 945 to align the various sampling module 948 portions and lancet module portions 950 allows the user to use a single lancet 183 of a particular lancet module portion 950 with multiple sampling module portions 948 of the sampling cartridge body 947. In addition, multiple different lancets 183 of lancet module portions 950 could be used to obtain a sample in a single sampling module portion 948 of the sampling cartridge body 947 if a fresh unused lancet 183 is required or desired for each lancing action and previous lancing cycles have been unsuccessful in obtaining a usable sample.

FIG. 102 shows an exploded view in perspective of the cartridge 945, which has a proximal end portion 954 and a distal end portion 955. The lancet cartridge body 946 is disposed at the proximal end portion 954 of the cartridge 945 and has a plurality of lancet module portions 950, such as the lancet module portion 950 shown in FIG. 103. Each lancet

module portion 950 has a lancet channel 951 with a lancet 183 slidably disposed within the lancet channel 951. The lancet channels 951 are substantially parallel to the longitudinal axis 952 of the lancet cartridge body 946. The lancets 183 shown have a drive head 198, shaft portion 201 and sharpened tip 5196. The drive head 198 of the lancets are configured to couple to a drive coupler not shown), such as the drive coupler 185 discussed above.

The lancets 183 are free to slide in the respective lancet channels 951 and are nominally disposed with the sharpened tip 196 withdrawn into the lancet channel 951 to protect the tip 196 and allow relative rotational motion between the lancet cartridge body 946 and the sampling cartridge body 947 as shown by arrow 956 and arrow 957 in FIG. 102. The radial center of each lancet channel 951 is disposed a fixed, known 15 radial distance from the longitudinal axis 952 of the lancetcartridge body 946 and a longitudinal axis 958 of the cartridge 945. By disposing each lancet channel 951 a fixed known radial distance from the longitudinal axes 952 and 958 of the lancet cartridge body 946 and cartridge 945, the lancet chan-20 nels 951 can then be readily and repeatably aligned in a functional arrangement with lancet channels 953 of the sampling cartridge body 947. The lancet cartridge body 946 rotates about a removable pivot shaft 959 which has a longitudinal axis 960 that is coaxial with the longitudinal axes 952 25 and 950 of the lancet cartridge body 946 and cartridge 945.

The sampling cartridge body 947 is disposed at the distal end portion 955 of the cartridge and has a plurality of sampling module portions 948 disposed radially about the longitudinal axis 949 of the sampling cartridge body 947. The longitudinal axis 949 of the sampling cartridge body 947 is coaxial with the longitudinal axes 952, 958 and 960 of the lancet cartridge body 946, cartridge 945 and pivot shaft 959. The sampling cartridge body 947 may also rotate about the pivot shaft 959. In order to achieve precise relative motion 35 between the lancet cartridge body 946 and the sampling cartridge body 947, one or both of the cartridge bodies 946 and 947 must be rotatable about the pivot shaft 959, however, it is not necessary for both to be rotatable about the pivot shaft 959, that is, one of the cartridge bodies 946 and 947 may be 40 secured, permanently or removably, to the pivot shaft 959.

The sampling cartridge body 947 includes a base 961 and a cover sheet 962 that covers a proximal surface 963 of the base forming a fluid tight seal. Each sampling module portion 948 of the sampling cartridge body 947, such as the sampling module portion 948 shown in FIG. 104 (without the cover sheet for clarity of illustration), has a sample reservoir 964 and a lancet channel 953. The sample reservoir 964 has a vent 965 at an outward radial end that allows the sample reservoir 964 to readily fill with a fluid sample. The sample reservoir 964 is in fluid communication with the respective lancet channel 953 which extends substantially parallel to the longitudinal axis 949 of the sampling cartridge body 947. The lancet channel 953 is disposed at the inward radial end of the sample reservoir 964.

The lancet channels 953 of the sample cartridge body 947 allow passage of the lancet 183 and also function as a sample flow channel 966 extending from an inlet port 967 of the lancet channel 953, shown in FIG. 106, to the sample reservoir 964. Note that a proximal surface 968 of the cover sheet 60 962 is spatially separated from a distal surface 969 of the lancet cartridge body 946 at the lancet channel site in order to prevent any fluid sample from being drawn by capillary action into the lancet channels 951 of the lancet cartridge body 946. The spatial separation of the proximal surface 968 of the 65 cover sheet 962 from the distal surface 969 of the lancet cartridge body 946 is achieved with a boss 970 between the

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two surfaces 968 and 969 that is formed into the distal surface 969 of the lancet cartridge body as shown in FIG. 105.

The sample reservoirs **964** of the sampling cartridge body 947 may include any of the sample detection analyte sensors, testing analyte sensors, analyte sensor contacts or the like discussed above with regard to other sampling module embodiments. The cover sheet 962 may be formed of PMMA and have conductors, analyte sensors or analyte sensor contacts formed on a surface thereof. It may also be desirable to have the cover sheet 962 made from a transparent or translucent material in order to use optical sensing or testing methods for samples obtained in the sample reservoirs. In the embodiment shown, the outer radial location of at least a portion of the sample reservoirs 964 of the sampling cartridge body 967 is beyond an outer radial dimension of the lancet cartridge body 946. Thus, an optical detector or analyte sensor 971, such as shown in FIG. 105, can detect or test a sample disposed within a sample reservoir 964 by transmitting an optical signal through the cover sheet 962 and receiving an optical signal from the sample.

The cartridge bodies 946 and 947 may have features, dimensions or materials that are the same as, or similar to, features, dimensions or materials of the sampling cartridges and lancet cartridges, or any components thereof, discussed above. The module portions 948 and 950 may also have features, dimensions or materials that are the same as, or similar to, features, dimensions or materials of the lancet or sampling modules, or any components thereof, discussed above. In addition, the cartridge 945 can be coupled to, or positioned adjacent any of the drivers discussed above, or any other suitable driver, in an operative configuration whereby the lancets of the lancet cartridge body can be selectively driven in a lancing cycle. Although the embodiment shown in FIGS. 102-106 allows for alignment of various sampling module portions 948 and lancet module portions 950 with relative rotational movement, other embodiments that function similarly are also contemplated. For example, lancet module portions, sampling module portions or both, could be arranged in a two dimensional array with relative x-y motion being used to align the module portions in a functional arrangement. Such relative x-y motion could be accomplished with position analyte sensors and servo motors in such an alternative embodiment order to achieve the align-

Referring to FIGS. 107 through 110, in one embodiment of the present invention, the disposable 1010 includes a plurality of penetrating members 1012 (lancets) and a plurality of analyte sensors 1014. The penetrating members 1012 are in a penetrating member housing 1016 and the analyte sensors 1014 are in an analyte sensor housing 1018. The analyte sensor housing 1018 is positioned in a surrounding relationship to the penetrating member housing 1016. The analyte sensor housing 1018 can be anchored to the penetrating member housing 1016 by a variety of means including but not limited to, posts and heat staking, locked at the ends using a pin and socket design, and the like.

The analyte sensor housing 1018 is fixed to the penetrating member housing 1016 by a variety of means, including but not limited to, heat staking with a post on the penetrating member housing 1016 extending through a hole in the analyte sensor housing 1018, gluing the two together, ultrasonically welding, laser welding, and the like.

In one embodiment, the analyte sensor 1014 is an electrochemical glucose analyte sensor 1014, that includes three electrodes and associated chemistry that can be in the form of a test strip. The test strip can be based on chrono-amperometry measurement techniques using glucose oxidase (Gox)

enzyme and N,N,N,N'-Tetramethyl-p-phenylenediamine (TMPD), as electron transfer mediators. The analyte sensor 1014 can be a screen-printed three-electrode system. The conducting layers can be made with a commercially available carbon paste. The reference and the counter electrodes can be made of Ag/AgCl. The working electrode can be made from the same commercial carbon paste blended with Gox and a the mediator. The composition of the working electrode material can be modified to lower the response time. A hydrophilic membrane can be provided with a surfactant that stabilizes an 10 otherwise sublimable mediator.

In this embodiment, each penetrating member 1012 is kept sterile, and each analyte sensor 1014 is kept dry. It is possible to lance and capture blood and transfer it to the analyte sensor 1014 in a single step. The penetrating member 1012 is aligned 15 with the analyte sensor 1014. The alignment provides that upon penetration by a penetrating member 1012 at a selected tissue site, blood flows into the analyte sensor 1014. This embodiment provides an integrated lancing and sample capture device with analyte sensing.

With the separate and integrated penetrating member housing 1016 and analyte sensor housing 1018, there is a separation of sterilizing the penetrating members 1012 and maintaining the analyte sensors 1014 dry.

In one embodiment, the disposable 1010 includes a plural- 25 ity of tests which may include used and unused penetrating members 1012, and used and unused analyte sensors 1014. The disposable 1010 can include micro fluidic configurations for transporting blood to the sensing area of the analyte sensors **1014** from a wound created in the skin by the penetrating member 1012. In one specific embodiment, the disposable 1010 has a penetrating member disk of fifty radially arranged penetrating members 1012 on a plastic disk, attached to which is the analyte sensor housing 1018, more particularly illustrated in FIGS. 111-114, which has fifty individual ana- 35 lyte sensors 1014, fifty individual desiccant pieces 1020 and fifty seals 1022.

In this embodiment, the analyte sensor housing 1018 is a circular ring which is attached to the penetrating member housing 1016. The analyte sensors 1014 are positioned radi- 40 front and rear seals 1022, associated with the analyte sensor ally outwardly relative to the penetrating members 1012. The analyte sensor housing 1018 can be molded as a circular ring, formed as a strip that is then curled or bent into a circle, and the like.

In one embodiment, the analyte sensors 1014 are cut or 45 punched from sheets into ribbons and then into individual analyte sensors 1014 which are inserted into the analyte sensor housing 1018, and then locked in place. The locking can be achieved by a variety of means including but not limited to, the use of glue, the use of ultrasonic welding, forming an 50 interference fit with rib structures 1024 that become deformed by the analyte sensors during fixation, and with the use of features that are formed after the analyte sensors 1014 are placed in the analyte sensor housing 1018, with at least portions of the features becoming deformed to trap the ana- 55 lyte sensors 1014 and fix them in place.

The analyte sensors 1014 are aligned to the penetrating members 1012. The penetrating members 1012 are aligned to the penetrating member housing 1016. The penetrating member housing 1016 is aligned to the analyte sensor housing 60 1018. The analyte sensors 1014 are aligned to the analyte sensor housing 1018. In one embodiment, the penetrating members 1012 are aligned to the penetrating member housing 1016 via bearings, the bottom part of the bearing being formed during a molding process of the penetrating member 65 housing 1016. The top of the bearings are formed after molding and after the penetrating members 1012 are inserted into

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the penetrating member housing 1016. The penetrating member housing 1016 is aligned to the analyte sensor housing 1018 via two posts that are formed in the penetrating member housing 1016, along with a mating hole and slot in the analyte sensor housing 1018. There are a variety of way for aligning the analyte sensor 1014 to the analyte member housing 1018. The edges of analyte sensors 1014 can be aligned with walls of the pockets 1019, which can house the analyte sensor 1014 and the desiccant 1020, that are formed in the analyte sensor housing 1018. Female trenches can be molded in the analyte sensors 1014 which align with mating male features in the analyte sensor housing 1018. Holes can also be formed in the analyte sensors 1014 which align with posts in the analyte sensor housing 1018.

The front and rear surfaces of the analyte sensor housing 1018 can have surfaces that are suitable for heat-sealing one or more vapor barrier films or seals, collectively denoted as 1022, including but not limited to aluminum foil, such that each analyte sensor 1014 is individually protected from the 20 environment. Each analyte sensor 1014 can then be held in its own, individual pocket 1019 in the analyte sensor housing 1018. In one embodiment, each pocket 1019 has its own desiccant 1020 so that each analyte sensor 1014 resides in a dry environment protected from the outside humidity as well as from adjacent analyte sensors 1014.

In one embodiment, the desiccant 1020 is co-molded during the injection molding of the analyte sensor housing 1018. The seals 1022 can be removed by a variety of different means, including but not limited to a punch. In one embodiment, one or more seals 1022 cover the pockets 1019 of the analyte sensor housing 1018 to prevent moisture ingress. The seals 1022 can be heat sealed while the analyte sensor housing 1018 is in a flat position and then bent to become attached to the penetrating member housing 1016. In another embodiment, the seals 1022 can be heat sealed in a rotary process around an analyte sensor housing 1018. In another embodiment the seals 1022 are sealed to the analyte sensor housing 1018 during a molding process.

In one embodiment, before a analyte sensor 1014 is used, 1014, are removed. After the seals 1022 are removed, a penetrating member 1012 can be actuated through or next to an opening 1021 in an associated analyte sensor support and enter the finger.

In one specific embodiment, the removal and/or opening of the seals 1022, such as by punching and the like, uses two separate punches. The first punch opens the penetrating member shaft compartment for gripping the penetrating member 1012 for actuation, opens the front compartment so the penetrating member can exit the sterilized penetrating member housing 1016 and also opens the back of the sealed analyte sensor housing 1018 to provide that the penetrating member 1012 can transect the analyte sensor support. A second punch opens the front compartment of the analyte sensor housing 1018 so that the penetrating member 1012 can exit the analyte sensor housing and create a wound in the skin.

In one embodiment of the present invention a body fluid sampling system 1010 is provided for use on a tissue site. A drive force generator is coupled to penetrating members 1012. The penetrating members 1012 are at least partially housed in a penetrating member housing 1016. The penetrating member housing 1016 has openings 1021 for the penetrating members 1012. The penetrating members 1012 are launched along a launch path by the drive force generator through an opening 1021 in the penetrating member housing 1016. Analyte sensors 1014 are housed in a analyte sensor housing 1018. Each analyte sensor 1014 is associated with a

penetrating member 1012. The analyte sensor housing 1018 and the penetrating member housing 1016 are coupled to provide that each analyte sensor 1014 is positioned out of a launch path of a penetrating member 1012. The analyte sensor housing 1018 is coupled with the penetrating member 5 housing 1016 to provide for a selected alignment of penetrating members 1012 to analyte sensors 1014. In one embodiment, the analyte sensor housing 1018 is locked to the penetrating member housing 1016.

The analyte sensor housing 1016 includes desiccant pock- 10 ets 1019 for receiving desiccant pieces/plugs 1020. Each desiccant plug 1020 is associated with a analyte sensor 1014. The desiccant plugs 1020 maintain the analyte sensors in a dry environment prior to launch of a penetrating member 1012 into a tissue site.

A first seal 1022 is in a sealing arrangement to at least tip portions of the penetrating members 1012 to provide that the tip portions remain in a sterile environment prior to launch of a penetrating member 1012 to a tissue site. In one embodiment, the first seal 1022 includes a plurality of seal portions 20 that are each associated with a separate penetrating member 1012. In another embodiment, a plurality of first seals 1022 are provided, each associated with a separate penetrating member 1012.

permits the penetrating members 1012 to be in a sterile environment. At least a third seal 1022 covers the desiccant pockets 1019.

The foregoing description of embodiments of the present invention has been presented for purposes of illustration and 30 description. It is not intended to be exhaustive or to limit the invention to the precise forms disclosed. Obviously, many modifications and variations will be apparent to practitioners skilled in this art. It is intended that the scope of the invention be defined by the following claims and their equivalents.

What is claimed is:

- 1. A body fluid sampling system for use on a tissue site, the system comprising:
  - a drive force generator;
  - in a penetrating member housing, the penetrating member housing having a plurality of openings with each of the plurality of openings associated with one of the plurality of penetrating members, each of the plurality of penetrating members being configured to be coupled to 45 housing has a ring shaped geometry. the drive force generator and be launched along a launch path through an opening in the penetrating member housing; and
  - a plurality of analyte sensors housed in an analyte sensor housing, each of an analyte sensor being associated with 50 one of the plurality of penetrating members, the analyte sensor housing and the penetrating member housing being coupled to provide that each of the plurality of analyte sensors is aligned to and positioned out of the bers,
  - one or more seals that seal the plurality of penetrating members in the penetrating member housing and the plurality of analyte sensors in the analyte sensor housing members sterile and each of the plurality of analyte sensors dry wherein the seals are removable for launch of one of the penetrating members and for use of one of the analyte sensors.
- 2. The system of claim 1, wherein the analyte sensor hous- 65 ing including a plurality of desiccant pockets each with a desiccant plug.

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- 3. The system of claim 2, wherein the desiccant plug is associated with one of the plurality of analyte sensors to keep it dry.
- 4. The system of claim 2, wherein a post on either of the penetrating member housing or the analyte sensor housing glues the two together.
  - 5. The system of claim 1, further comprising:
  - a first seal in the sealing arrangement to at least tip portions of the penetrating members to provide that the tip portions remain in a sterile environment prior to launch of one of the plurality of penetrating members to a tissue
- 6. The system of claim 5, wherein the first seal includes a 15 plurality of seal portions, each of the plurality of seal portions associated with one of the plurality of penetrating members.
  - 7. The system of claim 5, wherein a plurality of first seals are provided, each of the plurality of first seals being associated with one of the plurality of penetrating members.
  - 8. The system of claim 1, wherein the desiccant plug maintains one of the plurality of analyte sensors in a dry environment about prior to launch of one of the plurality of penetrating members into a tissue site.
- 9. The system of claim 1, wherein the analyte sensor hous-In another embodiment, at least a second seal 1130 that 25 ing is coupled with the penetrating member to provide for a selected alignment of penetrating members to analyte sen-
  - 10. The system of claim 1, wherein the analyte sensor housing is configured to lock to the penetrating member housing.
    - 11. The system of claim 5, further comprising:
    - at least a second seal that permits the penetrating members to be in a sterile environment and the desiccant pockets in a dry environment until launch of one of the plurality of penetrating members.
    - 12. The system of claim 11, further comprising: at least a third seal that covers the desiccant pockets.
- 13. The system of claim 1, wherein each of the plurality of a plurality of penetrating members at least partially housed 40 analyte sensors has its own housing pocket, a front seal and a back seal, wherein each housing pocket is a desiccant pocket.
  - 14. The system of claim 1, wherein the penetrating member housing has a disc shaped geometry.
  - 15. The system of claim 1, wherein the analyte sensor
  - **16**. The system of claim **1**, wherein the analyte sensor is positioned for blood flow into the analyte sensor upon launch of one of the plurality of penetrating members and penetration of the penetrating member at a skin surface.
  - 17. The system of claim 1, wherein the analyte sensor housing is heat staked to the penetrating member housing.
  - 18. The system of claim 1, wherein the analyte sensor housing is glued to the penetrating member housing.
  - **19**. The system of claim **1**, wherein the analyte sensor launch path of one of the plurality of penetrating mem- 55 housing is ultrasonically welded to the penetrating member housing.
    - 20. The system of claim 1, wherein the analyte sensor housing is laser welded to the penetrating member housing.
  - 21. The system of claim 1, wherein each of the plurality of separately to keep each of the plurality of penetrating 60 analyte sensors is fixed in the analyte sensor housing with glue.
    - 22. The system of claim 1, wherein each of the plurality of analyte sensors is fixed in the analyte sensor housing by ultrasonic welding.
    - 23. The system of claim 1, wherein each of the plurality of analyte sensors is fixed in the analyte sensor housing with an interference fit.

- 24. The system of claim 1, wherein rib structures are used to retain each of the plurality of analyte sensors, each of an analyte sensor at least partially deforming associated ribs to create an interference fit.
- 25. The system of claim 1, wherein features are used to fix 5 each of the plurality of analyte sensors, the features being formed after each of an analyte sensor is placed in the analyte sensor housing.
- **26**. The system of claim **1**, wherein bearings align the penetrating members to the penetrating member housing.
- 27. The system of claim 1, wherein the penetrating member housing is aligned to the analyte sensor housing with posts that are formed in the penetrating member housing, and a mating hole and slot in the analyte sensor housing.
- **28**. The system of claim **1**, wherein edges of analyte sensors are aligned with walls of pockets formed in the analyte sensor housing.
- **29**. The system of claim **1**, wherein female trenches are molded in the analyte sensors and align with mating male features in the analyte sensor housing or male trenches are 20 molded in the analyte sensors and align with mating female features in the analyte sensor housing.
- **30**. The system of claim **1**, wherein holes formed in the analyte sensors align with posts in the analyte sensor housing.

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